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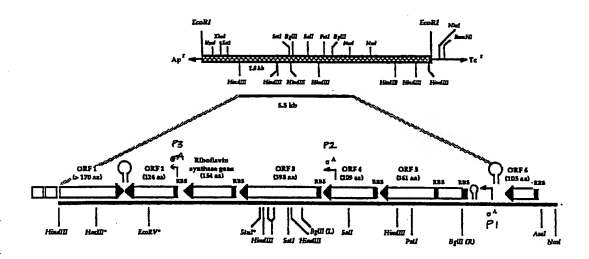
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(A) Riboflavinoverproducing strains of bacteria.

The present invention is directed to bacteria that overproduce riboflavin. The invention concerns the nucleotide sequence of the rib operon and its open reading frames, and recombinant bacteria that contain the rib operon. Specifically, the invention relates to bacteria that have been mutated so that their production of riboflavin and/or purines is deregulated, and which have copies of the rib operon inserted and amplified in their chromosomal DNA. In a specific embodiment, the rib operon itself can be deregulated by replacing its control regions with sequences that allow constitutive or unregulated expression. The bacteria, operons and sequences of this invention are used to overproduce relatively large amounts of riboflavin by fermentation.

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FIGURE 4.



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RIBOFLAVINOVERPRODUCING STRAINS OF BACTERIA

Riboflavin (vitamin B₂) is synthesized by all plants and many microorganisms but is not produced by higher animals. Because it is a precursor to coenzymes such as flavin adenine dinucleotide and flavin mononucleotide, that are required in the enzymatic oxidation of carbohydrates, riboflavin is essential to basic metabolism. In higher animals, insufficient riboflavin can cause loss of hair, inflammation of the skin, vision deterioration, and growth failure.

Riboflavin can be commercially produced either by a complete chemical synthesis, starting with ribose, or by fermentation with the fungi Eremothecium ashbyli or Ashbya gossypii (The Merck Index, Windholz et al., eds., Merck & Co., p. 1183, 1983). Mutants of Bacillus subtilis, selected by exposure to the purine analogs azaguanine and azaxanthine, have been reported to produce riboflavin in recoverable amounts (U.S. Patent No. 3,900,368, Enel et al., 1975). In general, exposure to purine or riboflavin analogs selects for deregulated mutants that exhibit increased riboflavin biosynthesis, because the mutations allow the microorganism to "compete out" the analog by increased production (Matsui et al., Agric. Biol. Chem. 46:2003, 1982). A purine-requiring mutant of Saccharomyces cerevisiae that produces riboflavin has also been reported (U.S. Patent No. 4,794,081, Kawai et al., 1988). Rabinovich et al. (Genetika 14:1696 (1978)) report that the riboflavin operon (rlb operon) of B. subtilis is contained within a 7 megadalton (Md) EcoRI fragment (later referred to as a 6.3 Md fragment in Chikindas et al., Mol. Genet. Mik. Virusol. no. 2:20 (1987)). It is reported that amplification of the rib operon may have been achieved in E. coli by cloning the operon into a plasmid that conferred resistance to ampicillin and exposing bacteria containing that plasmid to increasing amounts of the antiblotic. The only evidence for rib amplification is a coincident increase in the presence of a green-fluorescing substance in the medium; the authors present a number or alternative possibilities besides an actual amplification of the operon to explain the phenomenon observed.

French Patent Application No. 2,546,907, by Stepanov et al. (published December 7, 1984), discloses a method for producing riboflavin that utilizes a mutant strain of B. subtilis which has been exposed to azaguanine and roseoflavin and that is transformed with a plasmid containing a copy of the rib operon.

Morozov et al. (Mol. Genet. Mik. Virusol. no. 7:42 (1984)) describe the mapping of the B. subtilis rib operon by assaying the ability of cloned B. subtilis rib fragments to complement E. coli riboflavin auxotrophs or to marker-rescue B. subtilis riboflavin auxotrophs. Based on the known functions of the E. coli rib genes, the following model was proposed for the B. subtilis operon: ribG (encoding a deaminase) - ribO (the control element) - ribB (a synthetase) - ribF - ribA (a GTP-cyclohydrolase) - ribT/D (a reductase and an isomerase, respectively) - ribH (a synthetase).

Morozov et al. (Mol. Genet. Mik. Virusol. no. 11:11 (1984)) describe the use of plasmids containing the B subtilis rib operon with either wild-type (ribO *) or constitutive (ribO 335) operator regions to assay their ability to complement B subtilis riboflavin auxotrophs. From the results, a revised model of the rib operon was proposed, with ribO now located upstream of all of the structural genes, including ribG, and with the existence of an additional operator hypothesized, possibly located just upstream of ribA.

Morozov et al. (Mol. Genet. Mik. Virusol. no. 12:14 (1985)) report that the B. subtilis rib operon contains a total of three different promoters (in addition to a fourth "promoter" that is only active in E. coll). The primary promoter of the operon was reported to be located within the ribO region, with the two secondary promoters reported between the ribB and ribF genes and within the region of the ribTD and ribH genes, respectively.

Chikindas et al. (Mol. Genet. Mlk. Virusol. no. 2:20 (1987)) propose a restriction enzyme map for a 6.3 Md DNA fragment that contains the rib operon of B. subtilis. Sites are indicated for the enzymes EcoRl, Pstl, Sall, EcoRV, Pvull and Hindill.

Chikindas et al. (Mol. Genet. Mik. Virusol. no. 4:22 (1987) report that all of the structural genes of the B. subtilis rib operon are located on a 2.8 Md Bglll-HindIII fragment and that the Bglll site is located between the primary promoter of the operon and the ribosomal-binding site of its first structural gene. As described infra, Applicants show that this Bglll site is actually located within the most-5 open reading frame of the rib operon, so that the 2.8 Md fragment described does not contain all of the rib structural genes. Thus, in contrast to the report of Chikindas et al., the 1.3 Md Bglll fragment does not contain the ribosomal-binding site of the first structural gene; insertions at this site lead to a riboflavin-negative phenotype. Consequently, any attempt to use this Bglll site to engineer the rlb operon in order to increase expression, for example by r placing the 5 regulatory region with a strong r promoter, would actually destroy the integrity of the first structural gene and thus the operon as well.

Chikindas t al. (Dokl. Akad. Nauk. 5 SSSR 298 :997 (1988)) disclose another model of the B. subtilis rib operon, containing the primary promoter, p₁, and twee minor promoters, p₂ and p₃: ribO(p₁)-ribG-ribB-p₂-



ribF-ribA-ribT-ribD-p₃-ribH As before, it is incorrectly reported that the 1.3 Md Bglll fragment contains the entire first structural gene of the operon and that this proximal Bglll site maps within the primary regulatory region.

The present invention is directed to bacteria that overproduce riboflavin. The invention relates to the nucleotide sequence of the rib operon and its open reading frames, and recombinant bacteria that contain the rib operon. Specifically, the invention is directed to bacteria that have been mutated so that their production of riboflavin and/or purines is deregulated, and to bacteria which have copies of the rib operon inserted and amplified within their chromosomal DNA. In one embodiment, the rib operon itself can be deregulated by replacing its control regions with sequences that allow constitutive or unregulated expression. The bacteria, operons and sequences of this invention can be used to produce large amounts of riboflavin by fermentation.

The present invention is illustrated by way of specific examples detailed below, for example, a mutant of B. subtilis 1A382, RB50::[pRF8]60(Ade*), is produced that is deregulated for riboflavin and purine production and has the rib operon amplified within its chromosome. This mutant is able to produce greater than 5 g/l of riboflavin after 48 hours of fermentation in a 14-liter vessel. Other bacteria are described in which riboflavin production is increased to over 10 g/l under similar conditions.

This invention generally features the production of large quantities (over 10 g/l) of riboflavin by construction of various bacterial strains and growth of those bacterial strains within a medium and under conditions suitable for production of the riboflavin. In a first aspect, the invention features a recombinant bacterium which includes at least one copy of an exogenously introduced nucleic acid within its chromosome. This nucleic acid encodes one or more riboflavin biosynthetic proteins, is heritable, and capable of expression by the bacterium such that riboflavin biosynthesis by the bacterium is increased relative to a bacterium lacking such a sequence.

By "recombinant bacterium" is meant a bacterium which contains one or more nucleic acid sequences, from the same or another organism, at a site at which those sequences do not naturally occur, and/or in a copy number in which they do not naturally occur. Thus, the term includes those bacteria in which two copies of a nucleic acid sequence, e.g., a gene or an operon, are provided at a site which normally includes only one copy of the sequence. It also includes bacteria in which one or more copies of a nucleic acid sequence are introduced at a site which does not normally include that sequence. Such recombinant bacteria are constructed by standard recombinant DNA technology.

By "exogenously introduced" is meant that the nucleic acid is introduced into the chromosome from a source outside of that chromosome by any standard technique, including recombinant DNA technology, transformation, and transfection. It also includes the progeny of such bacteria, for example, those bacteria produced by cellular division of an originally constructed, transformed, or transfected bacterium.

By "riboflavin biosynthetic proteins" is meant to include those peptides, polypeptides or proteins which are directly involved in the synthesis of riboflavin from guanosine triphosphate. These proteins may be identical to those which naturally occur within a bacterium and are involved in the synthesis of riboflavin within that bacterium. Alternatively, they may be modifications of such proteins, for example, they may contain modifications which do not significantly affect the biological activity of the protein. For example, the natural protein may be modified by introducing or substituting one or more amino acids, preferably by conservative amino acid substitution, or by removing nonessential regions of the protein. Such modifications are readily performed by standard techniques.

In some embodiments, the bacterium contains two or more copies of the nucleic acid sequence; and the nucleic acid encoding one or more of the riboflavin biosynthetic proteins is present at at least two sites within the chromosome of the bacterium.

By "site" is meant a distinct chromosomal location relative to a wild-type bacterium at which the nucleic acid encoding the biosynthetic proteins is located. For example, such nucleic acid may be located at the naturally occurring site for genes encoding such proteins (i.e., at a rib locus), or it may be located at a site distant from this location. Preferably such distant sites are chosen from regions of chromosomal nucleic acid which are not essential to the recombinant bacterium, such as regions which encode proteins which are not essential to production of riboflavin. Examples of such regions include those which encode certain extracellular enzymes such as proteases. Insertion at such sites does not interfere with a desirable quality or trait. Any site is suitable as long as the functioning of the bacterium, with regard to riboflavin production, is not substantially affected.

In other embodiments, the nucleic acid is present in a plurality of copies at one or more of the sites; and the nucleic acid is present at at least three sites within the chromosome. By introducing the nucleic acid at different sites, the total number of copies of the nucleic acid within the chromosome can be increased. Increasing the copy number, increases the amount of riboflavin pr duction.

Generally the riboflavin biosynthetic proteins are encoded by one or more rib genes (e.g., an inactivation of which creates a riboflavin auxotroph), preferably at least five distinct rib genes identifiable from the nucleotide sequence provided in Fig. 3. Preferably, at least five copies of such genes are provided. By "rib genes" is meant those genes or portions of genes which encode proteins which occur naturally within an organism, or perform a similar function to such proteins, which are involved in the biosynthetic conversion of guanosine triphosphate to riboflavin within a bacterium.

In a related aspect, the invention features a recombinant bacterium which includes nucleic acid encoding one or more riboflavin blosynthetic proteins, e.g., the gene products identified as ORF1 and ORF6 in Fig. 4, the expression of at least one of which is controlled by a transcription element not naturally associated with the nucleic acid. Alternatively, the recombinant bacterium includes one or more rib genes or transcription units the expression of which is controlled by a transcription element not naturally associated with that rib gene.

By "transcription element" is meant to include any nucleic acid which effects (i.e., turns on) the transcription of nucleic acid downstream from that transcription element. Examples of such elements include promoters and operators. Such transcription elements are not naturally associated with the nucleic acid, for example, they may be heterologous transcription elements. That is, they may be isolated from other species or genera of bacteria or other organisms. Alternatively, the transcription element may be one naturally present in the bacterium but not normally associated with a rib gene to which it is now transcriptionally linked. Such elements do not include those which are naturally associated with a rib gene.

In other embodiments, the recombinant bacterium includes at least three (or at least five) rib genes and the expression of all three rib genes is controlled by a transcription element not naturally associated with those rib genes; at least two transcription elements are provided; the rib genes are provided within the chromosome of the recombinant bacterium; the recombinant bacterium is deregulated for riboflavin gene expression; and the transcription element is a promoter. For example, the promoter is a constitutive, growth-regulated, or inducible promoter, such as one associated with the SPO1 phage, and/or veg, amy, and sacQ-sensitive promoters, e.g., apr.

By "deregulated" is meant that the level of riboflavin production is greater than that observed in a bacterium with natural riboflavin regulatory systems (i.e., a wild type bacterium). Examples of such deregulated bacteria include those which are resistant to various purine analogs or antagonists whereby such analogs or antagonists can e.g. be selected from the group consisting of 8-azaguanine, psicofuranine, decoyinine, 8-azaguanine, sulfaguanine, 6-thioguanine, and methionine sulfoxide, and/or riboflavin analogs e.g. roseoflavin. Preferred deregulated bacteria are resistant to at least one of the following purine analogs or antagonists 8-azaguanine or decoyinine and roseoflavin.

In other specific embodiments, at least one of the rib genes includes a ribosome binding site not naturally associated with the rib gene; the rib genes are present at two sites within the chromosome; and the rib genes are present in a plurality of copies within the chromosome. In more preferred embodiments, the rib genes are Bacillus rib genes, for example ORF3 and ORF4 shown in Fig. 4, and the transcription element is located in a region 5'-upstream of ORF3 or ORF5; and the rib genes are chosen from a β -riboflavin synthase-encoding gene, ORF2, ORF3, ORF4, and ORF5; and the bacterium belongs to a species of Escherichia, e.g., E. coli, Bacillus, e.g., B. subtilis, Klebsiella, or Cornyebacterium.

In another related aspect, the invention features nucleic acid, which includes five or more rib genes, the expression of which is controlled by a transcription element not naturally associated with that rib gene.

In a further aspect, the invention relates to a vector comprising first a nucleic acid sequence of bacterial especially Bacillus whereby B. subtilis is preferred or E. coli or yeast origin coding for one or more, whereby at least five is preferred, riboflavin biosynthetic proteins and second one or more whereby one or two is preferred transcription elements not naturally associated with this nucleic acid sequence. Such vectors are preferred whereby the transcription elements are as already specified above. Further preferred are vectors as specified in the Examples e.g. pRF50, pRF69 pRF70, pRF71, pRF78, pRF81 and/or pRF89.

Another aspect of the invention relates to a recombinant bacterium comprising a bacterium which has been transformed by a vector as specified above whereby at least one copy, preferably a plurality of copies of said nucleic acid sequence including said transcription elements has been introduced at one or more sites with in its chromosome and said nucleic acid sequence including said transcription elements is heritable and capable of expression by the bacterium, such that riboflavin biosynthesis by the bacterium is increased relative to a bacterium lacking such nucleic acid sequence including said transcription elements. Introduction at one or two such sites is preferred. Furthermore recombinant bacteria are preferred whereby the bacterium which is transformed is already deregulated for riboflavin gene expr ssion. Of such deregulated bacteria E. coli or Bacillus specially B. subtilis strains are preferred whereby B. subtilis strains RB50 and RB58 are especially pref rred.



The invention relates also to a process for the preparation of a recombinant bacterium whereby a bacterium especially as already specified above is transformed by a v ctor comprising a nucleic acid sequence of bacterial or yeast origin coding for one or more riboflavin biosynthetic proteins or a vector as already specified above whereby at least one copy, preferably a plurality of copies of said nucleic acid sequence optionally including said transcription elements is introduced at one or more, preferably one or two sites with in its chromosome and said nucleic acid sequence optionally including said transcription elements is heritable and capable of expression by the bacterium, such that riboflavin biosynthesis by the bacterium is increased relative to a bacterium lacking such nucleic acid sequence optionally including said transcription elements.

The invention relates also to bacteria as specified herein wherein one or more transcription elements naturally associated with the rib biosynthetic genes in the chromosome of such bacteria have been replaced by transcription elements as specified. Such replacements can be made on the basis of the description of the present invention and by methods known in the art.

In another aspect, the invention features a method or process for production of riboflavin. The method includes growing cells especially recombinant bacteria as specifically mentioned herein under suitable growth conditions. Such suitable growth conditions are characterized by limiting the availability of a component of the growth medium and/or feed medium in such a way that aerobic conditions for the growth of said recombinant bacterium are maintained. Such conditions can be also characterized e.g. by maintaining a level of dissolved oxygen at a concentration between about 5% to 30%. The man skilled in the art is familiar with the fact that such levels of dissolved oxygen can vary dependent on the specific technical equipment used for growing said recombinant bacteria and for measuring said dissolved oxygen concentration. Under anaerobic conditions the synthesis of riboflavin is reduced. In some embodiments, the limiting component is chosen from a carbon source, nitrogen source, or a component required by the cells (e.g., in the feed medium). For example, if the cells are auxotrophic, for example, for methlonine, a limiting level of methionine may be provided in the growth medium. In another example, such component could be glucose or a carbonic acid, e.g. a citric acid cycle acid, such as citric acid or succinic acid, or an amino acid.

In a related aspect, the invention features another method for increasing production of riboflavin by a bacterium. In this method, the strain of bacterium used is deregulated for riboflavin production. More than one copy of a nucleic sequence encoding one or more riboflavin biosynthetic proteins is introduced into the chromosomal DNA of this bacterium. Preferably the bacterium used in this method is selected from one of those described above. The invention relates also to a method for the production of riboflavin by growing a recombinant bacterium obtained by a process as specified above under conditions as also specified above.

In other aspects of the invention, purified nucleic acid and the recombinant polypeptide product of such nucleic acid is provided. Generally, the purified nucleic acid consists essentially of all or a portion of the rib operon, for example, the specific open reading frames shown in Fig. 3. Such purified nucleic acid may be provided within a vector such as a plasmid, phage, or cosmid, or may be integrated within the chromosome of a bacterium. This nucleic acid is separated from nucleic acid with which it is naturally linked. For example, 6.5 kb of the nucleic acid encoding the whole rib operon may be inserted within a Bacillus subtilis chromosome at a site distant from that site in which the 6.5 kb DNA is normally present. By recombinant polypeptide is meant biologically active protein free of extraneous polypeptide (i.e., not fused to a heterologous polypeptide) having an enzymatic activity equivalent to such a naturally produced polypeptide.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

The drawings will first briefly be described:

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Figure 1. The riboflavin biosynthetic pathway, modified from Keller et al., Blochem. 27:1117 (1988). The corresponding intermediates shown are those produced by E. coll (which are presumably the same as those produced by B. subtilis): structure 1, guanosine triphosphate (GTP); structure 2, 2,5-diamino-6-(ribosylamino)-4(3H)-pyrimidinone-5'-phosphate; structure 3, 5-amino-6-(ribosylamino)-2,4(1H,3H)-pyrimidinedione-5'-phosphate; structure 4, 5-amino-6-(ribitylamino)-2,4(1H,3H)-pyrimidinedione-5'-phosphate; structure 5, 6,7-dimethyl-8-ribityllumazine; structure 6, riboflavin. The biosynthetic enzymes indicated are those encoded by B. subtil is (GTP cyclohydrolase, α and β subunits of riboflavin synthase) or those proposed to be encoded by B. subtilis (a rib-specific deaminase, and a rib-specific reductase). Figure 2. Schematic representation of purine biosynthesis. The purine biosynthetic pathway, including the portion responsible for riboflavin biosynthesis, is depicted. The individual enzymes of the pathway are identified by their gene symbols (E. coli nom nclature). Abbreviations are as foll ws: PRPP, phosphoribosylpyrophosphate; GAR, glycinamide ribonucleotide; pur, GAR formyltransf rase; PRA, phosphoribosylamine; purA, adenylosuccinate synthetase; purB, adenylosuccinate synthetase; FGAR,

formylglycinamide ribonucleotide; SAICAR, aminoimidazolesuccinocarboxamide ribonucleotide; purC, SAICAR synthetase; FGAM, formylglycinamidine ribonucleotide; purD, GAR synthetase; AIR, aminoimidazole ribonucleotide; purE, AIR carboxylase; CAIR, carboxyaminoimidazole ribonucleotide; purF, PRPP amidotransferase; AICAR, aminoimidazolecarboxamide ribonucleotide; purH, AICAR formyltransferase; purJ, inosine monophosphate (IMP) cyclohydrolase; FAICAR, formamidoimidazolecarboxamide ribonucleotide; purL, FGAR amidotransferase; guaA, guanosine monophosphate (GMP) synthetase; purM, AIR synthetase; guaB, IMP dehydrogenase.

Figure 3. The complete nucleotide and deduced amino acid sequences of the B. subtilis rib operon. The nucleotide sequence was determined by dideoxy sequencing of M13 clones. The deduced amino acid sequence is indicated by the one letter code (Lehninger, Biochemistry. 2d Ed., Worth Publishers, Inc., New York, p. 72).

Figure 4. A schematic representation of the rib gene cluster. The top diagram is the restriction endonuclease map of the cloned 10 kb EcoRl DNA fragment in plasmid pRF2, containing the B. subtilis rib operon. Homology to the 54-mer probe specific for the riboflavin synthase gene is depicted by the solid bar. The hatched box depicts Rib cloned DNA, while the thin black line represents pBR322 DNA. The bottom diagram is based on the complete nucleotide sequence of the 6.0 kb fragment to which the rib operon was localized. Open reading frames are depicted by open boxes, with arrows indicating the direction of transcription, and closed boxes indicating the putative ribosome binding sites. Probable σ^A promoter regions are shown. Tentatively identified rho-independent transcription termination sites are indicated by a "hairpin" symbol. Not all restiction sites are indicated.

Figure 5. Strain lineage of RB50. The lineage of the riboflavin overproducing strain of B. subtilis, RB50, is depicted. The various parent strains were exposed to riboflavin and purine analogs to select appropriate mutations.

Figure 6. Origins of rib*s recombinant plasmids. A schematic diagram of the production of the rib operon-containing recombinant plasmids pRF1, pRF2, pRF3, pRF6 and pRF7 is presented. A library of size-selected, 9-11 kb fragments of B. subtilis DNA was used to produce a gene library in E. coli plasmid vectors. Clones were selected by hybridization to the 54-mer probe specific for the β subunit of the riboflavin synthase gene.

Figure 7. The strain lineage of B. subtilis RB53::[pRFB]₉₀. Plasmid pRFB was integrated into the chromosome of the intermediate strain RB52 and amplified; the resulting strain was exposed to the purine analog azaguanine.

Figure 8. Identification of regions essential for riboflavin biosynthesis using insertions and deletions. A diagram is presented of the 10 kb cloned EcoRl DNA fragment with the regions essential for riboflavin biosynthesis indicated. Insertions and deletions at the indicated restriction sites enabled the localization of the rib operon. Not all restriction sites are indicated.

Figure 9. Hairpin-loop structures of the possible rho-independent transcription termination sites. Their locations in the nucleotide sequence of Figure 3 are shown below each structure. Also presented are their free energies of formation, determined according to Tinoco et al. (Nature (London) New Biology 246 :40 (1973)).

Figure 10. Structure of various plasmid derivatives used in S-30 in vitro coupled transcription translation reactions. A schematic diagram is shown of the rib operon regions contained in the plasmid derivatives used in the S-30 reactions, as well as the open reading frames predicted to be expressed.

Figure 11. Comparison of riboflavin production curves. Riboflavin production curves for various fermentation protocols are shown. Open squares: RBF-14 using RB50::[pRF8]₆₀ (Ade⁻). Closed squares: RBF-22 using RB50::[pRF8]₆₀ (Ade⁻). Closed circles: RBF-29 using RB50::[pRF8]₆₀ (Ade⁻).

Figure 12. Construction of pRF40.

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Figure 13. Construction of pRF50.

Figures 14, 15, and 16. Structure of various vectors.

Figure 17, 55-mer used in plasmid construction.

Figure 18. Various oligonucleotides used in vector construction.

In the practice of the present invention, host bacterial strains are derived that contain one or more mutations in genes of the riboflavin biosynthetic pathway or in the biosynthetic pathway of various purines, which mutations lead to riboflavin overproduction. In one embodiment, such mutations lead to riboflavin overproduction by deregulating steps in the riboflavin biosynthetic pathway. In another embodiment, the mutations increase riboflavin production by causing an inhibition in the use in an alternative metabolic pathway of a precursor for riboflavin biosynthesis.

In a specific embodiment, desired mutations in the genetic background of the host bacteria can be



induced by exposure to analogs of purine or riboflavin that compete with their authentic counterparts in the metabolic pathways of the host; bacteria that surviv such exposure will hav mutations that allow them to overproduce the authentic counterpart to the analog, thus "competing out" the purine or riboflavin analog that would otherwise be lethal. The biosynthesis of riboflavin in B. subtilis originates with guanosine triphosphate (Figure 1, structure 1). Guanosine triphosphate (GTP), via guanosine monophosphate (GMP), is a product of the purine biosynthetic pathway (Figure 2). In a preferred embodiment, to obtain a host strain that overproduces riboflavin, one can use classical genetics to both increase the amount of GTP that the cell produces and to deregulate the riboflavin pathway. Purine overproduction in B. subtilis can be achieved by obtaining mutants resistant to purine analogs or antagonists. Examples of some of the purine analogs that can be used include but are not limited to 8-azaguanine (Ishii and Shiio, Agric. Biol. Chem. 36:1511, 1972; Konishi and Shiro, Agric. Biol. Chem. 32:396, 1968), psicofuranine and decoyinine (Matsui et al., Agric. Biol. Chem. 43:1739, 1979; Matsui et al., Agric. Biol. Chem. 43:393, 1979), 8-azaxanthine, sulfaguanine, 6-thioguanine (Debabov, V.G. In The Molecular Biology of the Bacilli vol. 1 Bacillus subtilis, D.A. Dubnau, ed. (Academic Press, New York), pp. 331-370, 1982) and others, and/or the antagonist methionine sulfoxide (Matsui et al., App. Env. Microbiol. 34:337, 1977), and any combination thereof.

The riboflavin pathway can be deregulated by obtaining mutants resistant to a riboflavin analog. An example of a riboflavin analog that can be utilized is roseoflavin (Matsul et al., Agric. Biol. Chem. 46:2003, 1982).

In a specific embodiment of the present Invention, bacteria that are mutationally resistant to the analogs azaguanine, decoyinine and roseoflavin, can be used. Specific mutants resistant to each of these compounds are described below. Bacteria with mutations rendering them resistant to other analogs can also be used. It is also deemed within the scope of the present invention to utilize bacteria with different mutations rendering resistance to these same analogs, or different combinations of these mutations, either in combination, with or without, various mutations to other analogs.

If exposure to the analog alone does not produce resistant mutants at a high enough frequency, various mutagens can be used to increase the frequency of mutation in general and thus increase the number of analog-resistant mutants. As one example, ethyl methyl sulfonate can be used, but other mutagens including but not limited to nitrosoguanidine or UV irradiation can also be used.

Suitable bacterial hosts include all Bacilli species (including in a preferred embodiment B. subtilis), E. coli, and many other gram-positive and gram-negative bacteria. Species which can recognize the promoter sequences of the cloned rib operon to be inserted within their genome are suitable for use. The plasmids described below can be used to introduce rib genes into other bacteria by standard procedure, e.g., transformation. Expression of the inserted rib genes can be determined by spectroscopy as described below, or by observation of the bacteria under UV light, as described below.

In addition to creating mutations by exposure to purine or riboflavin analogs, bacterial strains that already contain mutations that are known to affect their purine or riboflavin biosynthetic pathways can be utilized. For example, the present invention makes use of but is not limited to B. subtilis strain 1A382, which contains the mutation pur-60, making it auxotrophic for adenine. Because this mutation blocks the utilization of the riboflavin precursor inosine monophosphate (IMP) in a metabolic pathway other than riboflavin production, increased amounts of IMP are available for riboflavin biosynthesis, thus increasing riboflavin production. There are many other mutations which can be utilized to potentially increase riboflavin production, including but not limited to guaC3, his—and others that are included within the scope of the present invention. The guaC3 mutation prevents the conversion of GMP back into IMP (see Figure 2), thus increasing the amount of riboflavin biosynthetic precursors available.

Suitable mutations affecting riboflavin overproduction can be mapped by various methods known in the art. In a specific embodiment, a mutation can be mapped by complementation of auxotrophic mutants.

The riboflavin biosynthetic genes from various bacteria can be cloned for use in the present invention. Yeast or bacterial cells from species including but not limited to the genus Bacilius, E. coli and many other gram-positive-and-gram-negative bacteria can potentially serve as the nucleic acid source for the molecular cloning of the rib operon. The DNA containing the rib operon may be obtained, by standard procedures known in the art, for example, from a DNA library prepared by cloning chromosomal DNA or fragments thereof, purified from the desired bacterial cell, into a suitable vector for propagation of the gene. (See, for example, Maniatis et al., 1982, Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1982, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K., Vol. I, II).

In the molecular cloning of the gene from chromosomal DNA, fragments are generated, some of which will encode the d sired rib peron. The DNA may b cleaved at specific sites using various restriction nzymes. Alternatively, one may use DNASe in the presence of manganese to fragment the DNA, or the

DNA can be physically sheared, as for example, by sonication. The linear DNA fragments can then be separated according to size by standard techniques, including but not limited to agarose and polyacrylamide gel electrophoresis and density gradient centrifugation.

Once the DNA fragments are generated, DNA librarles are prepared using an appropriate cloning and/or expression vector. A large number of vector-host systems known in the art may be used. Possible vectors include, but are not limited to, plasmids or modified viruses, but the vector system must be compatible with the host cell used. For E. coli such vectors include, but are not limited to, bacteriophages such as λ derivatives, high-copy plasmids such as pBR322 or pUC plasmids, or low-copy plasmids derived from Pseudomonas plasmid RK2. For Bacillus such vectors include, but are not limited to, bacteriophages such as ρ11 Dean et al., J. Virol. 20: 339, 1976; Kawamura et al., Gene 5:87, 1979) or Δ105 derivatives (lijima et al., Gene 9:115, 1980; Errington, J. Gen. Microbiology 130:2615, 1984; Dhaese et al., Gene 32: 181, 1984; Errington, J. in Bacillus Molecular Biology and Biotechnology Applications, A.T. Ganesan and J.A. Hoch, eds. (Academic Press, New York,), p. 217, 1986), high-copy plasmids such as pUB110 (Ehrlich, Proc. Natl. Acad. Sci. (USA) 74: 1680, 1977) or PBD64, or low-copy plasmids such as pE194 derivatives (Gryczan, T.J. in The Molecular Biology of the Bacilli, D.A. Dubnau, ed. (Academic Press, New York), pp. 307-329, 1982; Horinouchi and Weisblum, J. Bacteriol. 150: 804, 1982). Recombinant molecules can be introduced into host cells via transformation, transfection, protoplasting, infection, electroporation, etc.

Once the DNA libraries are generated, identification of the specific clones harboring recombinant DNA containing the rib operon may be accomplished in a number of ways (as described, for example, in Maniatis et. al., supra). For example, if an amount of the operon or a fragment thereof is available from another bacterial source (e.g., from E. coli) and is sufficiently homologous to the riboflavin biosynthetic genes of Bacillus to hybridize thereto, that DNA can be purified and labeled, and the generated bank of DNA fragments may be screened by nucleic acid hybridization to the labeled probes (Benton, W. and Davis, R., 1977, Science 196:180; Grunstein, M. and Hogness, D., 1975, Proc. Natl. Acad. Sci. U.S.A. 72:3961). Alternatively, sequences comprising open reading frames of the endogenous rib operon, or subsequences thereof comprising about 10, preferably 15 or more nucleotides, may be used as hybridization probes. Such probes can be made synthetically, based on a portion of the nucleic acid or amino acid sequence (examples of which are provided below) of a gene product known to be encoded by the operon ("reverse genetics"). If a purified rib operon-specific probe is unavailable, cloned gene libraries of restriction fragments (from partial Sau3A-digests, for example) can be made in bacteria, especially B. subtilis or E. coli, and the rib operon-containing recombinant clones can be identified by either marker-rescue or complementation of known rib mutations.

In a preferred embodiment, the <u>rib</u> operon of B. subtilis can be isolated for use from an E. coli plasmid library of B. subtilis DNA. In particular, and as described below, the B. subtilis rib operon can be isolated by virtue of its homology to a radiolabelled, synthesized nucleotide probe that is derived from an internal region of a gene product known to be encoded by the operon of B. subtilis. Although a portion of the amino acid sequence for β-riboflavin synthase (Ludwig et al., J. Biol. Chem. 262:1016, 1987) can be the basis for such a probe, with the third nucleotide of each codon estimated from frequency of codon usage, a similar probe based on another region of this protein or another protein from the rib operon can be utilized and would fall within the scope of the present invention. The present invention further enables screening by use of synthetic probes which are derived from the nucleic acid sequence shown in Fig. 3.

Analogous methods to those detailed here can be used to isolate the rib operon of other bacteria, especially other Bacilli or E. coli. In a specific embodiment, such clones can be selected by assay for ability to hybridize to the labeled B. subtilis rib operon or a hybridizable portion thereof. It is well known in the art that starting from an appropriate mRNA preparation, cDNA can be prepared; such cDNA can also be used in accordance with the present invention to prepare vectors for the transformation of appropriate bacteria for riboflavin overproduction.

Once the host cells with recombinant DNA molecules that include the isolated rib operon or a portion thereof are identified, the DNA may be obtained in large quantities. This then permits the rib operon to be manipulated and its nucleotide sequence to be determined using various cloning and sequencing techniques familiar to those knowledgeable in the art.

For example, insertional mutagenesis can be used to locate and characterize the rib operon and genes thereof within a cloned piece of DNA. In a specific embodiment, rib-biosynthetic containing regions can be identified by inserting small cat (chloramphenicol acetyltransferase)-containing restriction fragments into several different restriction enzyme sites of the cloned DNA, and testing each derivative for insertional inactivation of riboflavin biosynthesis in an appropriate host (see below).

The cloned DNA corresponding to the rib operon can be analyzed by methods including but not ilmited to Southern hybridization (Southern, E.M., 1975, J. Mol. Biol. 98:503-517), Northern hybridization (see ...9.,

Freeman et al., 1983, Proc. Natl. Acad. Sci. U.S.A. 80: 4094-4098), restriction endonuclease mapping (Maniatis et al., 1982, Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York), and DNA sequence analysis. Restriction endonuclease mapping can be used to roughly determine the genetic structure of rib operon. Restriction maps derived by restriction endonuclease cleavage can be confirmed by DNA sequence analysis.

DNA sequence analysis can be performed by any techniques known in the art, including but not limited to the method of Maxam and Gilbert (1980, Meth. Enzymol. 65:499-560), the Sanger dideoxy method (Sanger, F., et al., 1977, Proc. Natl. Acad. Sci. U.S.A. 74:5463), or use of an automated DNA sequenator (e.g., Applied Biosystems, Foster City, CA). As an example, the DNA sequence of the rib operon of B. subtilis is presented in Figure 3.

Once the nucleotide sequence of the rib operon has been determined, putative open reading frames (ORFs) can then be identified along with the deduced amino acid sequence of their encoded product. Actual identification of the encoded product can be carried out, e.g., by performing S-30 coupled in vitro transcription/translation reactions, with various ORFs used as templates. Various mutational derivatives of the ORFs can also be tested for activity in functional assays of the S-30 reaction products, in order to test the function of the encoded products.

In a specific embodiment of the invention relating to the B. subtilis rib operon, and detailed in the examples below, the above-described methods were used to determine that B. subtilis riboflavin biosynthesis is controlled by a single operon of approximately 4.2 kb containing five biosynthetic genes: the β subunit of riboflavin synthase and ORFs designated 2, 3, 4, and 5 (see Figure 4). ORFs 2, 3, 4, and 5 were subsequently shown to encode proteins with molecular weights of about 15 kd, 47 kd, 26 kd, and 44 kd, respectively. As described below, ORF 5 was shown to encode a putative rib-specific deaminase that catalyzes the reduction of a deaminated pyrimidine to a ribitylamino-linkage in an early step in riboflavin biosynthesis. Our data also indicated that ORF 4 encodes the α subunit of riboflavin synthase and ORF 3 encodes a GTP cyclohydrolase, while ORF 2 possibly encodes a rib-specific reductase. ORF 1 and ORF 6 were found to be outside the primary transcription unit of the rib operon. The primary site for initiation of transcription of the rib operon was determined to be probably the apparent σ^A promoter located 290 bp upstream from the first gene in the operon, ORF 5 (Figure 4, P₁). The coding regions, promoters and transcription termination sites of the B. subtilis rib operon are shown in Table VI below.

The present invention encompasses the nucleotide and amino acid sequences of the genes of the rib operon, as well as subsequences thereof encoding functionally active peptides, and sequences which are substantially the same as such sequences. A functionally active peptide, as used herein, shall mean a protein or peptide which is capable of catalysing a reaction leading to riboflavin biosynthesis. A functionally active nucleic acid sequence shall mean a sequence capable of regulating riboflavin biosynthesis. A sequence substantially the same as another sequence shall mean a sequence capable of hybridizing to the complementary sequence thereof. In addition, a nucleic acid sequence not naturally controlling the expression of a second nucleic acid sequence shall mean a sequence which does not control the expression of the second sequence in the bacterium from which the second sequence is isolated.

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Once the genetic structure of the rib operon is known, it is possible to manipulate the structure for optimal use in the present invention. For example, the rib operon can be engineered to maximize riboflavin production.

Depending on the host-vector system utilized, any one of a number of suitable transcription and translation elements may be used. Promoters produced by recombinant DNA or synthetic techniques may also be used to provide for transcription of the inserted sequences. When propagating in bacteria the regulatory sequences of the rib operon itself may be used. In an embodiment in which the entire rib operon, or greater than one gene thereof, is desired to be expressed as a polycistronic message, a prokaryotic host is required. In an embodiment in which a eukaryotic host is to be used, appropriate regulatory sequences (e.g., a promoter) must be placed in the recombinant DNA upstream of each gene/ORF that is desired to be expressed.

Specific Initiation signals are also required for efficient translation of inserted protein coding sequences. These signals include the initiation codon (ATG, GTG or TTG) and adjacent sequences, such as the ribosome binding site (RBS). It should be noted that the RBS of a given coding sequence can be manipulated to effect a more efficient expression of that coding sequence at the translational level. In cases where an entire open reading frame of the rib operon, including its own initiation codon and adjacent regulatory sequences, is inserted into the appropriate expression vectors, no additional translational control signals may be needed. However, in cases where only a portion of the coding sequence is inserted, or where the native regulatory signals are not recognized by the host cell, exogenous translational control signals, including the initiation codon, must be provided. The initiation codon must furthermore be in phase

with the reading fram of the protein coding sequences to ensure translation of the entire insert. These exogenous translational control signals and initiation codons can be of a variety of origins, both natural and synthetic.

In addition, a host cell strain may be chosen which modulates the expression of the rib operon gene(s) or modifies and processes the gene product(s) thereof in the specific fashion desired. Expression from certain promoters can be elevated in the presence of certain inducers; thus, expression of the genetically engineered rib operon proteins may be controlled. In one embodiment, the regulatory regions of the operon, such as the promoter and the termination/ anti-termination regulatory sequences, can be manipulated or replaced with constitutive or growth-regulated promoters to deregulate the rib operon and thus increase riboflavin production. Furthermore, appropriate cell lines or host systems can be chosen to ensure the desired modification and processing of the expressed proteins. Many manipulations are possible and within the scope of the present invention.

In one specific embodiment of the invention, the 5' regulatory sequence of the B. subtilis rib operon can be removed and replaced with one or more of several B. subtilis promoters; such a construction will cause high-level expression of the rib biosynthetic genes. This approach would involve the introduction of new restriction sites within a 20-30 bp region between the end of the transcription terminator and the RBS sequence of the first gene in the operon ORF 5. Such restriction sites can be introduced by either site-directed mutagenesis or by deleting all regulatory sequences upstream from the right-most BgIII (BgIII_R) site located within the first 30 bp of ORF 5 (see Figures 3 and 4) and inserting at this site a synthetic oligonucleotide that finishes off the 5' end of ORF 5 (including the ribosomal-binding site) and contains new upstream restriction sites. Once these constructions are made, promoter-containing restriction fragments with ends compatible to the new restriction sites can be introduced, causing expression of the rib genes under the control of the new promoter. Both constitutive and growth-regulated B. subtilis promoters can be used, including but not limited to strong promoters from the lytic bacteriophage SPO genes, veg, amy (amylase), and apr (subtilisin).

In another aspect of the invention, rib operon DNA fragments which have transcriptional regulatory activity (e.g., promoters) can be used to regulate the expression of heterologous gene products.

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According to the present invention, the rlb operon can be introduced into bacteria, including for example, Bacilli and E. coli, where it is expressed in a preferred embodiment, the bacterial host is one of the mutant hosts described above. In a specific embodiment, the cloned rib operon is integrated into the host chromosomal DNA, where it is replicated and expressed along with host genomic DNA. In a most preferred embodiment, multiple copies of the rib operon are integrated into the host chromosomal DNA, thus providing for amplified expression of the rib operon in the deregulated host. One method in which this may be accomplished is chromosomal insertion of a cat-containing rib operon followed by chloramphenicol amplification of the operon, as detailed in the examples sections infra. One can also use a tet^r gene, or certain other drug resistance genes that are expressed in Bacillus, with the same technique.

In specific embodiments, integration vectors containing the rib operon fragment can be engineered so as to contain the rib operon on the smallest possible DNA fragment, in an attempt to obtain greater amplification of the vector within the host chromosome. For example, vector DNA sequences may be deleted, and/or nonessential DNA flanking the rib operon can be deleted.

In general, bacteria that are prototrophic for riboflavin survive on minimal medium in the absence of riboflavin. Production of riboflavin can be detected and quantified by various methods. In a preferred embodiment, overproduction of riboflavin is readily observed when overproducing bacteria are exposed to UV light at 368 nm, as described infra, producing an observable, yellow fluorescence. For example, many of the engineered plasmids of the present invention are produced in E. coli. For some of these plasmids, overproduction of riboflavin has been confirmed by this method. The amount of riboflavin produced can be quantitated, e.g., with reverse-phase high performance liquid chromatography (HPLC). Cell-free supernatants from bacteria can be fractionated over an HPLC column, as described infra, and monitored for riboflavin at 254 nm. By extrapolation from a standard curve, the concentration of riboflavin can be determined by the area of the peak on the chromatogram.

Riboflavin can also be quantitated by fluorescence spectrophotometry. For example, samples containing riboflavin can be read in a fluorescence spectrophometer set at an emission wavelength of 525 nm and an excitation wavelength of 450 nm.

In addition, other methods known in the art are available to detect or quantitate riboflavin based on its physical and biological properties.

Riboflavin overproducing bacteria can be grown in vess is ranging from shake flasks to large "batch" fermentors, by methods known in the art (see below). In a preferred embodiment, nutrient feed can be manipulated to maximiz riboflavin production at the minimum cost by varying the nutrients in the medium.

In a specific embodiment, amplified <u>rib</u> -containing g nes can be maintained at high-copy number in the bacterial chromosome by the inclusion of about 60 µg/ml chloramphenicol in the inoculum seed strain (but not necessarily in the fermentor). Chemap 14-liter fermentors can be used at 1000 rpm with a head pressure of 0.6 atmospheres. To achieve a high cell density, various carbon sources such as, for example, glucose, sucrose, citric acid cycle acids, maltose or starch, and various nitrogen sources such as yeast extract, com steep liquor, ammonia and/or protein hydrolysates can be used. It is, however, essential to add parts of these media ingredients in such a way that limitations unfavorable to riboflavin production (e.g., oxygen starvation resulting from too high a carbon source concentration) are avoided. Fermentation media and conditions suitable for use are detailed below.

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Examples

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Example 1: Riboflavin-Overproducing B. subtilis Mutants

We describe in the examples herein the production of strains of Bacillus subtilis which overproduce riboflavin. In order to accomplish this, we used classical genetics, genetic engineering, and fermentation. Classical genetics with selection using purine and riboflavin analogs was used to deregulate the pathways for purine (riboflavin precursor) and riboflavin biosynthesis. Riboflavin production was increased further by cloning and engineering the genes of the riboflavin biosynthetic pathway (the rib operon), allowing for constitutive, high-level production of rate-limiting biosynthetic enzyme(s).

The biosynthesis of riboflavin in B. subtilis originates with GTP (Figure 1). To obtain a host that overproduces riboflavin we used classical genetics to both increase the amount of GTP that the cell produces and to deregulate the riboflavin pathway. Purine overproduction in B. subtilis can be achieved by obtaining mutants resistant to purine analogs such as azaguanine and decoyinine, and other antagonists such as methionine sulfoxide (see e.g., Ishii and Shiio, Agric. Biol. Chem. 36 (9):1511-1522, 1972; Matsui et al., Agric. Biol. Chem. 43 (8):1739-1744, 1979). The riboflavin pathway can be deregulated by obtaining mutants resistant to the riboflavin analog roseoflavin (Matsul et al., Agric. Biol. Chem. 46(8):2003-2008, 1982). Roseoflavin-resistant strains were selected from several strains which had been previously mutagenized and which were resistant to several purine analogs. Described below are the methods used to produce a strain (RB50) which overproduces riboflavin.

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8-Azaguanine-Resistant Mutants

B. subtilis is effectively killed by the purine analogue 8-azaguanine (Sigma Chemical Co., St. Louis, MO) at a concentration of 500 μg/ml, and resistant mutants appear spontaneously at a frequency of less than 1 in 108. Ethyl methyl sulfonate (EMS; Sigma) at 30 μg/ml was used as a mutagen to increase the frequency of azaguanine-resistant (Ag') mutations. Mutagenesis was performed on cells from B. subtilis strain 168 by standard procedures (Miller, 1972, Experiments in Molecular Genetics, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York). After plating 4 x 106 mutagenized cells on minimal medium (Sloma et al., J. Bact. 170:5557, 1988) containing 500 μg/ml azaguanine and restreaking for single colonies, 35 Ag' colonies resulted. One mutant, RB11 (Agr -11), was used in the construction of RB50.

Decoyinine-Resistant Mutants

Decoyinine-resistant (Dc') mutations were obtained spontaneously at a frequency of 1 in 10⁶ or after EMS mutagenesis at 1 in 10⁵ by plating cells on minimal medium containing 100 μg/ml of decoyinine (Upjohn Co., Kalamazoo, Ml). A Dc' mutant of RB11 was obtained by mutagenesis with EMS as described above. One Dc' colony, RB15 (Ag' -11, Dc' -15), was used in the construction of RB50.

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Transfer of the Ag and Dc Mutations

These purine analog-resistant mutations were transferred to a different strain background in order to

isolate th m from any unwanted EMS-induced mutations and to verify that th Agr and Dcr mutations were due to single loci. Since part of the "carbon flow" from inosine monophosphate (IMP), a riboflavin precursor, is also used for adenine nucleotide biosynthesis, a host strain was selected that was blocked in the adenosine monophosphate (AMP) pathway via the mutation pur-60, allowing more carbon material to "flow" from IMP to the guanine nucleotide precursors of riboflavin (Figure 2). B. subtilis strain 1A382 (hisH2, trpC2, pur-60) was made competent (Sloma et al., J. Bact. 170:5557 (1988)) and transformed (by the method of Gryczan et al., J. Bact. 134:318 (1978)) with total DNA prepared from the Agr/Dcr mutant RB15. The Trp*-(tryptophan) revertant colonies were selected, with 3.3% (10/300) of those also being Dcr and 2.3% (7/300) Agr. This result was not unexpected since, due to "congression" (transformation of a second unlinked marker), a number of the Trp* colonies should also be resistant to decoyinine or azaguanine.

One Dc' colony, RB36 (his H2, pur-60, Dc'-15), one Ag' colony, RB40 (his H2, pur-60, Ag'-11), and one Dc'/Ag' colony (which was also found to be his), RB39 (pur-60, Ag'-11, Dc'-15), were all selected for further study.

Methionine Sulfoxide-Resistant Mutants

Selection using high levels of methionine sulfoxide (MS; 10 mg/ml, Sigma) resulted in spontaneous mutants appearing at a sufficiently high frequency that mutagenesis with EMS was not necessary. The Agr/Dcr mutant, RB39, was streaked onto minimal medium containing 10 µg/ml MS. Resistant colonies were obtained and were restreaked for single resistant colonies. One strain, RB46 (pur-80, Agr-11, Dcr-15, MSr-46) was selected for further study.

5 Roseoflavin Resistant Mutants

Although many of these Agr, Dcr and MSr mutants were likely to be overproducing GTP, none of them produced levels of riboflavin detectable on plates. In order to deregulate the riboflavin biosynthetic pathway, conditions were determined to select for resistance to the riboflavin analog roseoflavin (Toronto Research Chemical). Maximum killing of cells occurred at 100 μ g/ml of roseoflavin in minimal or complete medium; increasing the concentration did not result in any additional killing. Mutations to roseoflavin resistance (RoF') spontaneously occurred at a sufficiently high rate (approximately 5 x 10⁻⁵) such that mutagenesis with EMS or other chemicals was not necessary.

Approximately 1000 RoF' colonies were obtained from each of the strains described above, 1A382, RB36, RB39, RB40 and RB46. RoF' mutants from all of these strains showed a low level of fluorescence on minimal media plates when exposed to long-wave UV light (366 nm), indicating some riboflavin production. One of the RoF' colonies obtained from RB46, RB46Y (pur-60, Ag'-11, Dc'-15, MS'-46, RoF'-46), when grown on minimal medium, produced 14 mg/l of riboflavin as determined by HPLC (described above).

Of all the strains treated, only RB39 and RB46 produced a significantly different phenotype when RoFr colonies were selected. Approximately 0.5% to 1.0% of the RoFr colonies of either RB39 or RB46 produced an intensely fluorescent, yellow colony. Of these colonies, RB51 (pu r -60, Agr-11, Dcr-15, RoFr-51), arising from RB39, and RB50 (pur-60, Agr-11, Dcr-15, MSr-46, RoFr-50), arising from RB46, produced a stable, fluorescent-yellow phenotype which correlated with a higher level of riboflavin production, as determined by HPLC. When grown in minimal medium, both RB50 and RB51 produced higher levels of riboflavin in their supernatants than the other RoFr strains, about 40 mg/l and 30 mg/l, respectively. The lineage of RB50 is depicted in Figure 5.

Because intensely fluorescent (and thus riboflavin overproducing) colonies could be obtained in non-MS' strains such as RB51, it appeared that this mutation in general might not be contributing significantly to the higher production phenotype. Both of the other mutations, Ag' and Dc' (Ag'-11 and Dc'-15 in RB39), appear to be necessary to produce high levels of riboflavin since no intensely fluorescent RoF' colonies could be found in strains containing only the Ag'-11 (from RB40) or Dc'-15 (from RB36) mutation alone.

guaC Mutations

Another possibly important mutation for achieving overproduction of GTP, and thus riboflavin, is guac3, which prevents the conversion of GMP back into IMP (see Figure 2). To construct a strain containing guac3 that overproduces riboflavin, competent B. subtilis strain 62121 cells (guaC3, trpC2, metC7) (Endo et al., J.

Bact. 15: 169, 1983) were transformed with RB50 DNA and selected for Dc^r on plates containing 100 μg/ml of decoyinine. Thousands of Dc^r colonies r sulted. Of 200 colonies which were patch d onto Dc^r plates, one was found that exhibited the riboflavin overproduction phenotype (based on UV fluorescence), and was RoF^r. This colony was designated RB52 (guaC3, trpC2, metC7, Dc^r-15, RoF^r-50) and was reserved for subsequent study.

Other Analog-Resistant Mutants

Finally, because mutants resistant to several additional purine analogs also have been reported to be altered in purine metabolism, such mutations were assayed in order to Investigate their effect on riboflavin-overproducing strains. It was determined that 500 g/ml of 8-azaxanthine, 1 mg/ml of 6-thioguanine, or 2 mg/ml of sulfaguanidine (Sigma) effectively kills wild-type B. subtilis. The azaguanine-resistant, riboflavin-overproducing strains RB50::[pRF8]₉₀ and RB53::[pRFB]₉₀ (see below) were found to be already resistant to azaxanthine. Although separate azaguanine-and azaxanthine-resistant mutations with different properties have been described previously, in this case the Ag'-11 and Ag'-53 mutations appear to also convey azaxanthine resistance.

20 HPLC Analysis of riboflavin in crude supernatants of B. subtilis

Accumulation of riboflavin in B. subtilis cultures was quantitated by reverse-phase HPLC. Riboflavin standards (Sigma Chemical Co., St. Louis, MO) or cell-free supernatants from strains to be tested were fractionated over a 4.6 mm x 250 mm Vydac C₁₈ column equilibrated with 1% ammonium acetate (pH 6.0). At injection, the column was developed with a linear gradient of methanol and monitored for riboflavin at 254 nm. Authentic riboflavin (i.e. riboflavin "standard") elutes at the mid-point of the gradient.

Example 2: Cloning B. subtilis rib operon

Our general strategy to isolate a restriction fragment containing the rib operon was to screen a "mini" E. coll plasmid library of B. subtilis DNA by hybridization with a synthetic oligonucleotide probe, the DNA sequence of which was partially derived from the published amino acid sequence for the β subunit of riboflavin synthase (Ludwig et al., J. Biol. Chem. 262:1016, 1987). A summary of the protocol is presented in Figure 6.

A synthetic, 54-base "guess-a-mer" oligonucleotide probe was used for this screening based on amino acids 84-102 of the 240 amino acid riboflavin synthase protein, sequenced by Ludwig et al. (J. Biol. Chem. 262:1016-1021, 1987). The third nucleotide of each codon in the probe was chosen according to estimates made of the most frequent codon usage of B. subtilis, based upon, for example, some of the sequences available in GenBank® (Los Alamos Nat. Lab, Los Alamos, NM). The probe consisted of the following sequence:

5'-GGAGCTACAACACATTATGATTATGTTTGCAATGAAGCTGCTAAAGGAATTGCT-3'.

To test the specificity of the probe, the ³²P-labelled 54-mer DNA was hybridized to nylon filters containing EcoRl-digested chromosomal DNA (Southern, J. Mol. Biol. 98:503, 1975) isolated from wild-type and the mutant B. subtilis strains. The probe strongly hybridized to a single 9-10 kb fragment of EcoRl-digested B. subtilis (ribs * met -) DNA, which is in good agreement with the predicted size of the rib-containing fragment (Osina et al., FEBS. Lett. 196:75, 1986). A labelled fragment of the identical size was detected when the probe was hybridized to two mutant strains, RB46 (pur-60, Ag'-11, Dc'-15, MS'-46) and RB50 (pur-60, Ag'-11, Dc'-15, MS'-46, RoF'-50), the latter being a riboflavin overproducer. These hybridization experiments were repeated using HindIII-cut chromosomal DNA, which resulted in the probe identifying a smaller, single fragment of approximately 1.8 kb; this latter result was useful in determining the general location of the rib biosynthetic operon within the cloned DNA.

55 Isolation of Plasmids pRF1, pRF2 and pRF3, Containing Wild-type rib Biosynthetic Genes

A "mini" gene library of 9-11 kb EcoRl fragm nts from B. subtilis strain 168 (rib*) DNA was prepared using pRK290, a low-copy number vector derived from the Pseudomonas replicon RK2 (Ditta et al., Plasmid

13:149, 1985). ECoRI fragments (size 9-11 kb) of B. subtilis (rib met) DNA were isolated by sucrose (10-40%) rate-zonal centrifugation. A four-fold excess of these fragments (0.22 μg) was ligated to EcoRI-cut pRK290 (0.26 μg), that had been dephosphorylated with calf intestinal alkaline phosphatase (CIAP), at a total DNA concentration of 10 μg/ml. Approximately 10 ng of ligated DNA was transformed into E. coli DH5 (F-, endAl, hsdR11 [r_k-, m_k+], supE₄₄, thi-1, λ-, recAl, gyrA96, relAl), resulting in tetracycline-resistant (Tc') colonies at a frequency of 7.7 x 10⁴/ μg of DNA. To determine the fraction of transformants containing insert DNA of 9-11 kb, plasmid mini-lysates were prepared from several Tc' transformants, and their DNA was analyzed by restriction enzyme digestion. About 40% of the Tc' transformants were found to contain single EcoRI-generated inserts of 9-11 kb.

Approximately 1140 of the Tc^r colonies were screened with the ³²P-labelled 54-mer probe specific for the riboflavin synthase gene. One colony gave a positive signal. Plasmid DNA, designated pRF1, was isolated from this clone and tested for Rib^{*}-marker rescue activity by transforming the DNA into B. subtilis 1A210 that contains the riboflavin-deficient mutation rib-2, and selecting for Rib^{*} prototrophic colonies. pRF1 transformed 1A210 to Rib^{*} prototrophy at a high frequency. Plasmid DNA from a randomly chosen Tc^r transformant falled to rescue this marker.

Restriction enzyme analysis revealed that pRF1 actually contained two EcoRI-fragment inserts, of 10 kb and 11 kb. To determine which fragment contained the rib operon, EcoRI-digested PRF1 was probed with the ³²P-labelled, 54-mer riboflavin synthase probe. The results indicated that only the smaller, 10 kb fragment cross-reacted with the probe. Moreover, when the 10 kb ECoRI fragment was recloned into the EcoRI site of pBR322, recombinant plasmids pRF2 and pRF3 resulted, representing the two possible orientations of insertion. Both plasmids were found to rescue the rib-2 mutation of B. subtilis 1A210 to prototrophy at a high frequency.

Isolation of Plamsids pRF6 and pRF7 Containing rib Biosynthetic Genes From RoFr-B. subtilis Strain RB50

RB50 is one of the RoF mutants of B. subtilis, produced as described above, that is deregulated for ribofiavin biosynthesis. It has been reported that approximately 80% of RoF mutations reside within the rib operon at the rib0 locus (Stepanov, et al., Genetika (USSR) 13:490, 1977). Like the wild-type rib operon, rib genes in RB50 were also contained on a 9-10 kb EcoRi fragment; thus this fragment was cloned using the protocol outlined in Figure 6, with pBR322 used as the cloning vector. Size-selected 9-11 kb EcoRi fragments (0.1 μg) from RB50 were prepared as before and ligated to a two-fold excess of ends of EcoRicut, dephosphorylated pBR322 DNA (0.34 μg) at a total DNA concentration of 22 μg/ml. Approximately 9 ng of ligated DNA was transformed into E. coli DHS, resulting in amplcillin-resistant (Ap¹) colonies at a frequency of 3.5 x 10⁵/μg of DNA.

Restriction enzyme analysis of plasmid DNA isolated from a sampling of 12 Apr colonies revealed that 50% contained plasmids with 9-11 kb EcoRI Inserts. Approximately 1140 Apr colonies were screened with the ³²P-labelled 54-mer probe specific for the riboflavin synthase gene by colony hybridization. Six colonies gave positive signals. Plasmids pRF6 and pRF7, isolated from two of these six colonies, were identified by restriction enzyme analysis as containing inserts with the same orientation as pRF2 and pRF3, respectively. In addition, both plasmids were able to marker-rescue the rib-2 mutation at high frequencies.

Example 3: Introducing rib DNA Into B. subtilis

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As described supra, the rib operon from both a wild-type strain and a RoFr mutant of B. subtilis were cloned as identical 10 kb EcoRl fragments into the EcoRl site of the E. coli replicon pBR322; the derivation of these recombinant plasmids is schematically diagrammed in Figure 6. To introduce the 10 kb EcoRl fragment containing the rib operon into B. subtilis in multiple copies, and thus further increase riboflavin production, we constructed a plasmid vector which would allow integration into the B. subtilis chromosome. The integrated DNA was amplified by selecting colonies that would grow at high drug concentrations of chloramphenicol.

Construction of and Transformation with Integrational rib Plasmids pRF4 and pRF8

To construct the integrational vector, the drug-resistance gene chloramphenicol acetyltransferase (cat), which is selectable in B. subtilis, was introduced into pRF2 and pRF6, the pBR322 vectors with the 10 kb

fragment from wild-type or RoF B. subtilis strains, respectively. The plasmids pRF2 and pRF6 were digested with BamHI, which cuts the plasmids uniquely within the pBR322 sequence, and dephosphorylated with CIAP. The cleaved DNA was ligated to a 1.3 kb BamHI fragment containing the cat gene (Youngman et al., Plasmid 12: 1-9, 1984), and the ligated DNAs then transformed into E. coli DHS cells (Hanahand, J. Mol. Biol. 166: 557, 1983). Approximately 80-90% of the Ap' transformants were chloramphenicol resistant (Cm'); restriction analysis of the isolated plasmids (Maniatis et al.) confirmed that plasmid DNA from the Cm' colonies contained the 1.3 kb fragment. The plasmid containing the wild-type riboflavin fragment and the cat gene was designated PRF4; the plasmid containing the cloned riboflavin fragment from the RoF' strain was called pRF8. (Since the RoF' mutation was subsequently shown to be outside the rib operon, these plasmids are presumably identical).

The plasmids pRF4 and pRF8 were transformed into four different B. subtilis strains: the riboflavin overproducer RB50 (Ag'-11, Dc'-15, MS'-46, RoF'-50), the RB50 parent RB46 (Ag'-11, Dc'-15, MS'-46), the RB50 parent 1A382 and IS75, a common laboratory strain. Competent IS75 and 1A382 cells were transformed with pRF4 or pRF8; these same plasmids were introduced into RB46 and RB50 by transformation of protoplasts (Chang and Cohen, Mol. Gen. Genet 168 :111-115, 1979). The pRF4 or pRF8 DNA integrated into each of these four strains was amplified by selecting for colonies that grew at higher chloramphenicol concentrations. In each strain, we were able to obtain colonies that grew in up to 60 μg/ml of chloramphenicol.

In addition, RB52 (guaC3, trpC2, metC7 Dcr-15, RoFr-50), produced by transforming the guaC3 B. subtilis strain 62121 with DNA from RB50, was made competent and transformed with pRF8. The integrated plasmid in one of the many Cmr colonies that resulted was amplified using 90 μg/ml of chloramphenicol. The resulting cells, RB52::[pRF8]₉₀, were grown to mid-log phase and plated on minimal media containing 500 μg/ml azaguanine. Approximately 20 Agr colonies resulted. One such colony seemed to produce a more intense fluorescence. The lineage of this strain, RB53::[pRF8]₉₀, is given in Figure 7.

Example 4: Riboflavin Overproduction by Strains Containing pRF4 or pRF8

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RB50 containing pRF4 or pRF8 displayed the riboflavin overproduction phenotype (yellow and UV-fluorescent colonies). Amplification of the rib DNA in a wild-type strain or the parent strains of RB50 did not yield yellow or UV-fluorescent colonies, a finding that indicates that the RoF mutation (which deregulates the biosynthesis of riboflavin) is required for chromosomal amplification of wild-type DNA to cause riboflavin overproduction. A series of shake flask fermentations were performed in 25 ml of riboflavin minimal medium (RMM, in Table I) in a 300 ml baffled flask (Bellco) to measure the production of riboflavin from RB50 that contained the integrated and amplified rib operon.

TABLE I

COMPOSITION C	COMPOSITION OF MEDIA		
RMM	<u>g/</u> 1		
Sodium glutamate	2.0		
Casamino acids (Difo	0.2		
KH₂PO₄	6.0		
K₂HPO₄ (NH₄)₂SO₄	14.0 2.0		
Sodium citrate	1.0		
MgSO ₄ .7H₂O Adenosine	0.2 0.05		
	pH 7.0 and autoclaved)		
Maltose	15.0		
(added as sterile 20% after autoclaving)	% solution		

The fermentations were run with strains RB46, RB50 and RB50 containing pRF4 amplified by selection for resistance to 30 μ g/ml of chloramphenicol (RB50::[pRF4]₃₀) and 90 μ g/ml of chloramphenicol (RB50::[pRF4]₃₀). At 24 and 48 hours, supernatant samples were removed and measured for riboflavin content by reverse-phase HPLC.

As shown in Table II, RB50::[pRF4]₃₀ produced 0.3 g/l of riboflavin, and RB50::[pRF4]₅₀ produced 0.7 g/l of riboflavin, in 48 hours, which is significantly more than that produced by the strains without rib amplification, such as RB46 and RB50.

TABLE II

QUANTITATIVE ANALYSIS OF RIBOFLAVIN-CONTAINING				
SUPERNATAN	TS FROM B. S	SUBTILIS		
Strain	Culture Time			
	(hours)	(g/l)		
RB46	24	0.009		
RB50	24	0.02		
RB50::[pRF4] ₃₀	24	0.1		
RB50::[pRF4] ₉₀	24	0.4		
RB46	48	0.007		
RB50	48	0.05		
RB50::[pRF4]30	48	0.3		
RB50::[pRF4] ₉₀	48	0.7		

^{*} Riboflavin was measured using an HPLC assay.

The dramatic increase in riboflavin production resulting from amplification of rib genes in the deregulated host argues that information encoded by the cloned DNA is rate-limiting for riboflavin biosynthesis.

Example 5: Mapping the RoFf-50 Mutation

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The RoF'-50 mutation in RB50 appeared to be critical to the riboflavin-overproduction phenotype. To identify and possibly move the mutation into different strain backgrounds it was necessary to map the location of the RoF'-50 mutation on the B. subtilis chromosome. Since pRF4 and pRF8 gave very similar levels of riboflavin production in all strain backgrounds, it seemed unlikely that the RoF'-50 mutation was located on the cloned 10 kb EcoRl, rib-containing fragment. More likely, the RoF'-50 mutation is an unlinked repressor-type mutation, possibly in ribC, a repressor mutation which has been reported to map in the lysaroD region of the B. subtilis chromosome (Chernik et al., Genetika (USSR) 15:1569, 1979). To determine whether the RoF'-50 mutation was linked or unlinked to the riboflavin operon, competent B. subtilis 1A210 (rib-2) cells were transformed with RB50 DNA, selecting for rib*. Thousands of rib* colonies resulted, and 200 colonies were patched onto tryptose blood agar base containing 100 g/ml of roseoflavin. No RoF' colonies resulted, and none of the colonies exhibited the riboflavin overproduction phenotype, confirming that the RoF'-50 mutation is not located in the rib operon.

Example 6: Locating rib Biosynthetic Genes Using CAT Insertional Mutagenesis

Figure 4 contains a restriction map of the rib-containing 10 kb EcoRl fragment of pRF2, prepared according to standard procedures. Restriction enzyme sites for Xbal, Bglll, Sstl, Hpal and Ncol are unique to the Insert DNA, whereas Sali and Pstl cut once in the insert and once in the vector, the insert does not contain any BaMHI, Xh I or Nhel restriction sites. Restriction enzyme HindIII cleaves the insert at multiple sit s; th 54-mer probe specific for the riboflavin synthase gene hybridized to an approximately 1.8 kb HindIII fragment, suggesting that the rib operon must also reside in the general area surrounding the Sall

and left-most Bgill (Bgill) sites.

In general, to d termine the boundari s of the rib operon, small cat-c ntaining restriction fragments were used to construct insertions and deletions in the rib -cloned DNA fragment of pRF2. E. coli plasmid pEccl served as the primary source of restriction fragments bearing a cat gene which confers chloramphenicol-resistance in both E. coli and B. subtilis. This plasmid, a derivative of pMI1101 (Youngman et al., Plasmid 12, 1-9, 1984) in which a non-essential region of the plasmid was removed by standard recombinant DNA techniques, contains a 1.3 kb cat-containing fragment flanked by the "polylinkers" of M13mp7, and therefore is capable of generating cat cassettes with either Smal, EcoRI, Sall or BamHi ends. To generate Sstl or Xval-ended fragments containing the cat gene, the 1.3 kb cat-containing BamHi fragment of pEccl was isolated, the ends modified with HindIII linkers, and the modified fragment cloned into the HindIII site within the polylinker region of pIC2OR, generating plasmid pEcc4.

Integrative plasmid derivatives were first constructed in E. coli and then transferred to the rib chromosomal locus of B. subtilis by DNA transformation. This was done by linearizing the plasmid by a restriction enzyme cut outside the cloned DNA insert, transforming competent B. subtilis strain 1A382 or PY79 (β^c , rib) cells with this cut DNA, and selecting for Cm'. Because the pBR322 replicon is unable to replicate in B. subtilis, and the cat gene is bounded on both sides by sequences homologous to the rib locus, the cat-containing insertion or deletion can only be inserted into the chromosome by a double-crossover recombination event to yield Cm' transformants. To determine whether the insertion or deletion inactivated riboflavin synthesis, Cm' colonies were assessed for growth on minimal medium agar plates with or without the presence of riboflavin (Rib phenotype).

As diagrammed in Figure 8, cat-containing restriction fragments were inserted by ligation into the individual restriction sites for Xbal, Sstl, Sall and Bgill of pRF2, inserted between the pair of Bgill or Ncol sites (generating deletions removing either a 2.0 kb Bgill fragment or a 0.8 kb Ncol fragment) or inserted into single Haelll and EcoRV sites of the approximately 1.8 kb Hindill fragment that hybridized to the ribspecific DNA probe, according to standard techniques. The results are shown in Table III.

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TABLE III

CHARACTERIZATION OF INSERTION AND DELETION DERIVATIVES OF rib± DNA

5	Insertion derivative	B. <u>subtilis</u> b <u>Riboflavin Phenotype</u>
	A(XbaI)	
	r	.
10	1	ND
,,	B(<u>Sst</u> I _L)	
•	r 1	+
	C(<u>Sat</u> l _R)	ND
15	r	
15	1	
	D(<u>Bgl</u> II _L)	
	r 1	· ———
	E(<u>Sal</u> I)	
20	r r	
	1 .	.
	F(<u>Bgl</u> II _R)	•
	r 1	
25	G(<u>Hae</u> III)	
	r	ND
	1	+
	H(<u>Eco</u> RV)	•
30	r 1	+ ND
	•	ND
	Deletion derivative	
35	Bgl	•
	<i>r</i> 1	
	Nco	
	r	+
40	1	+
	a uru (right) and ulu (last) ilantis. the
	t (Light) and 'i' (cation of the inserted <u>cat</u> gene
45	relative to the restr	iction map in Figure 8.
~~	•	
	b B. subtilis strain lA	382 (rib+, trpC2, pur-60,
	hisH2) or PY79 (SP B ^C , rib ⁺)	

As summarized in Figure 8 and Table III, insertions into the Sall, either BgllI, or the "right most" Sstl (Sstl_R) sites, or deletion of the 2.0 kb BgllI fragment, all generated Cm' colonies that could not produce riboflavin (Rib⁻), indicating that the rib operon was centrally located within the cloned DNA. Significantly, removal of the 0.8 kb Ncol fragment apparently had no effect on riboflavin production (Rib⁺), suggesting that one end of the rib gene cluster was located to the left of the "left most" Ncol (Ncol₁) site. The other end of the rib operon was initially determined to map within the approximately 1.8 kb HindIII fragment because the two insertions at sites within the fragment, EcoRV and HaeIII, as well as sites distal to the fragment, Xbal and Sstl_L, all generated Cm^r colonies that produced riboflavin.

Example 7: Nucleotide Sequence of the rib Operon

Based on the cat-insertional mutagenesis of the cloned 10 kb DNA fragment, the entire rib operon was localized within a 6.0 kb region bounded by the Sstl_L and Ncol_L sites.

This 6.0 kb region of pRF2 containing the rib operon and flanking regions was sequenced by the dideoxy method of Sanger et al. (Proc. Natl. Acad. Sci. USA 74:5463, 1977). Briefly, M13 clones for sequencing were prepared either by subcloning specific restriction fragments into M13, by using the exonuclease activity of T4 DNA polymerase to generate a series of overlapping deletions (Dale et al., Plasmid 13:31, 1985), or by "shot-gun" cloning random fragments, from sonicated restriction fragments, into M13. In some cases, the nucleotide sequence across a restriction site juncture of adjacent fragments was also determined by primer extension sequencing. Approximately 5500 bp were sequenced on both strands and analyzed for sequences resembling typical open reading frames with gram positive-bacteria ribosome binding sites, gram-positive promoters and rho-independent transcription termination sites.

Analysis revealed six complete, non-overlapping open reading frames (Figure 3): ORF 2 (124 amino acids), the gene coding for the \$\beta\$ subunit of riboflavin synthase (154 amino acids), ORF 3 (398 amino acids), ORF 4 (215 amino acids), ORF 5 (361 amino acids) and ORF 6 (105 amino acids). Each ORF was preceded by a strong Bacillus ribosome binding site (RBS) with calculated thermostability ranging from \$\Delta\$G = -16 to -22 kcal/mol, and all of them were oriented in the same transcriptional direction. In addition, within the coding region of ORF 5, a second RBS site and ATG start codon were identified, potentially encoding a smaller protein of 248 amino acids. However, based on S-30 in vitro coupled transcription/translation reactions (see below), ORF 5 appears to encode only a 361 amino acid protein. Finally, part of another coding region, ORF 1, encoding the last 170 amino acids of a protein and oriented in the opposite direction, was also identified.

Based on the following observations, riboflavin biosynthesis in Bacillus is controlled by a single operon containing 5 genes: the β riboflavin synthase gene, ORF 2, ORF 3, ORF 4, and ORF 5, of which at least four, the β -riboflavin synthase gene, ORF 3, ORF 4 and ORF 5, unambiguously encode biosynthetic enzymes, with the remaining one, ORF 2, possibly encoding a biosynthetic enzyme.

- 1. ORF 3, ORF 4 and ORF 5 overlap restriction enzyme sites where insertion of cat-containing restriction fragments caused inactivation of riboflavin production in B. subtilis (Figures 4 and 8).
- 2. ORF 1 overlaps a restriction enzyme site(s) where insertion of cat-containing restriction fragments did not cause inactivation of riboflavin production in a rib B. subtilis strain (Table III and Figure 8), nor did it cause reduction of riboflavin production in the deregulated, RoF B. subtilis strain RB52.
- 3. ORF 2 also overlaps a restriction enzyme site, EcoRV, where insertion of cat-containing restriction fragments did not cause inactivation of riboflavin production in a rib B. subtilis strain (Table III and Figure 8). However, such an insertion did cause a detectable reduction of riboflavin production in the deregulated, RoF B. subtilis strain RB52, indicating that the mutated ORF 2 gene product was partially inactive for riboflavin production. The results suggest that ORF 2 does encode a rib-specific enzyme.
- 4. Two DNA sequences capable of forming stem-loop structures indicative of rho-independent transcriptional termination sites were identified within the intercistronic gaps between ORF 1 and ORF 2 and between ORF 5 and ORF 6 (Figures 4 and 9). Removal of structures between ORF 5 and ORF6 enhances expression of riboflavin. The structures impart riboflavin sensitivity to lacZ-fusion constructs; thus, they can be used to impart such sensitivity to any other gene to which they are fused at the 5 end upstream of the promoter.
- 5. A DNA sequence, TTGCGT-(17bp)-TATAAT, resembling the promoter recognized by the $_{0}^{A}$ (vegetative form) of B. subtilis RNA polymerase was identified approximately 290 bp upsteam from ORF 5, oriented in the same transcriptional direction as ORF 5 (Figure 4). A transcriptional fusion of this promoter (P₁, on a 1.1 kb Bglli-Ncol restriction fragment) to a promoterless E. coli lacZ gene (P₁-lacZ) displayed riboflavin-regulated expression of β -galactosidase activity in a rib $^{+}$, B. subtilis strain (62121) and high-level, constitive (unregulated) expression of β -galactosidase activity in a rib $^{+}$, RoF B. subtilis strain (RB52) only when the promoter was oriented in the same transcriptional direction as the gene, as shown in Table IV. Primer extension analysis was used to confirm the start site. Transcriptional and Northern analyses were used to show a polycistronic RNA of 4.2 kb encompasses the entire rib operon.

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TABLE IV

RIBOFLAVIN-REGULATED EXPRESSION OF P1-LacZ TRANSCRIPTIONAL FUSIONS				
β-Galactosidase Specifi Activity (Miller Units)				
Strain (integrated plasmid)	+ Riboflavin (2 µg/ml)	- Riboflavin		
B . subtilis 62121 (P ₁ - lac Za)	1.3	4.2		
B . subtilis RB52 (P ₁ - lac Z ^a)	31	38		
B . subtilis 62121 (P ₁ - lac Z ^b)	<0.1	<0.1		
B . subtilis 62121	<0.1	<0.1		

^a P₁ and lac Z orientated in the same direction ^b P₁ and lac Z orientated in opposite directions Based on these results, this σ^A promoter, P₁, is a primary promoter for transcription of ORF 5, ORF 4, ORF 3, β-riboflavin synthase gene and ORF 2.

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- 6. A second DNA sequence, TTGAAG-(17bp)-TACTAT, resembling a promoter recognized by the σ^A -(vegetative form) of B. subtilis RNA polymerase was identified within the 3 end of ORF 4, approximately 295 bp upstream from ORF 3 and oriented in the same transcriptional direction as ORF 3 (Figure 4). Integration into B. subtilis by a Campbell-type recombination event of an E. coli plasmid containing this promoter sequence on a 0.7 kb Sall-BgIII restriction fragment did not cause inactivation of riboflavin production in B. subtilis, results which indicated that this second sequence (P₂) has promoter activity and thus may actually control transcription (in addition to the σ^A P₁ promoter) of ORF 3, the β subunit riboflavin synthase gene and ORF 2. LacZ fusions and Northern analysis confirmed the existence of this promoter.
- 7. A third DNA sequence, TTGAAT-(18bp)-TAAAAA, possibly resembling the promoter recognized by the σ^A (vegetative form) of B. subtilis RNA polymerase was identified within the intercistronic region between the β subunit of the riboflavin synthase gene and ORF 2, approximately 83 bp upstream of ORF 2 and oriented in the same transcriptional direction (Figure 4). This σ^A promoter, P₃, may also control transcription of ORF 2, in addition to P₁ and P₂.
- 8. In vitro-coupled transcription/translation analysis of S-30 reactions of the cloned DNA confirmed that ORF 2, ORF 3, ORF 4, and ORF 5 all actually encoded proteins of the size predicted from their respective sequences.
 - 9. Three of the five presumed enzymatic steps in riboflavin biosynthesis were assigned to specific coding regions by comparing predicted amino acid sequences or molecular weights of their products to published protein sequences, using GenBank®, or known protein sizes.
 - a. The putative protein encoded by the open reading frame between ORF 2 and ORF 3 almost identically matched the published 154 amino acid sequence of the β subunit for the riboflavin synthase enzyme (Ludwig et al., J. Biol. Chem. 262:1016, 1987). Only one amino acid difference was detected: lysine was substituted for glycine at residue 65. This enzyme is reported to catalyze the formation of 6.7-dimethyl-8-ribityllumazine from 5-amino-6-ribitylamino-2,4(1 H , 3 H)-pyrimidinedione-5'-phosphate (Figure 1, structures 5 and 4, respectively) and 3, 4-dihydroxybutanone-4-phosphate.
 - b. A 39% identity in an 88-amino acid overlap was identified between the putative product of ORF5 and deoxycytidylate deaminase, a 188 amino acid protein encoded by the E. coll bacteriophage T₂ (Maley et al., J. Biol. Chem. 258:8290, 1983). Based on this result, ORF 5 most likely encodes the rib-specific deaminase that catalyzes the formation of 5-amino-6-(ribosylamino)-2,4(1H,3H)-pyrimidinedione-5-phosphate from 2,5-diamino-6-(ribosylamino)-4(3H)-pyrimidinone-5-phosphate (Figur 1, structures 3 and 2, resp. ctively).

c. The predicted molecular weight of the ORF 4 gene product (26,000 Da) was in good agreement with the molecular weight of the α -subunit for riboflavin synthase (23,000 Da; Bacher et al., J. Biol. Chem. 255:632, 1980). Based on this result, ORF 4 encodes the α -subunit for riboflavin synthase, which catalyzes the final step of the biosynthetic pathway: dismutation of 6,7-dimethyl-8-ribityllumazine to riboflavin (Figure 1, structures 5 and 6, respectively) and 5-amino-6-ribitylamino-2,4-(1H, 3H)-pyrimidinedione.

10. The remaining enzymatic steps in riboflavin synthesis were tentatively assigned to coding regions by aligning the position of ORFs to a physical map of rib mutations in the operon (Morozov et al., Mol. Genet. Mik. Virusol. no. 7:42 (1984)). Mutations for defective GTP cyclohydrolase were reported to map to the 0.5 kb Hindlll fragment. Since ORF 3 encompasses this restriction fragment, we concluded that ORF 3, at least in part, encodes this enzymatic function, which catalyzes the formation of 2,5-diamino-6-(ribosylamino)-4(3H)-pyrimidinone-5'-phosphate from GTP (Figure 1, structures 2 and 1, respectively). In addition, the biosynthetic gene encoding a rib-specific reductase was reported to be contained entirely within the approximately 1.8 kb Hindlll fragment. Since this fragment contains only two complete coding regions, the β subunit of the riboflavin synthase gene and ORF 2, we speculate that ORF 2 encodes the reductase, which catalyzes the formation of 5-amino-6-(ribitylamino)-2,4(1H,3H)-pyrimidinedione-5'-phosphate from 5-amino-6-(ribosylamino)-2,4(1H,3H)-pyrimidinedione-5'-phosphate (Figure 1, structures 4 and 3, respectively).

In addition, a similar rho-independent transcription termination site was detected in the apparent leader region of the operon, downstream of the putative σ^A P_1 promoter but just upstream of the first coding region of the operon, ORF 5 (Figures 4 and 9). This potential terminator structure may be involved in regulation of the rib operon by a termination/ anti-termination mechanism. In addition, a roseoflavin-resistant (R_oF^R) dependent regulatory region is present on a 0.7 kb Sall-BgllI restriction fragment of ORF3.

Assignment of rib ORFs to Protein Products

One method for confirming whether the rib-specific ORFs encode proteins is to "visualize" the size and number of proteins synthesized from the cloned DNA in an S-30 in vitro coupled transcription/translation reaction using PRF2 and its various derivatives as templates. The S-30 fraction kit (New England Nuclear; used according to manufacturer's specifications) is especially efficient in translating B. subtilis genes due to the presence of their strong ribosome binding sites.

Using the cloned 10 kb EcoRI fragment of pRF2 or pRF4 as templates, we expected to detect five putative rib-specific proteins: β riboflavin synthase, 14.7 kilodaltons (kd) (Ludwig et al., J. Biol. Chem. 262:1016, 1987); and the proteins from ORF 2, 13.6 kd; ORF 3, 43.7 kd; ORF 4, 23 kd; and ORF 5, 39.7 kd. We also expected to detect at least two other proteins, encoded by ORF 6 (11.6 kd) and ORF 1 (at least 18.7 kd), as well as any additional proteins encoded by genes present in the unsequenced regions of the 10 kb cloned DNA fragment. In addition, vector-associated proteins, including the bla and cat antibiotic resistance gene products, were also expected (the tet gene is not strongly expressed in S-30 reactions).

Excluding the bla- and cat-specific proteins (32 kd and 18 kd, respectively) and other vector-associated proteins, a total of six major ³⁵S-labelled proteins were detected, with molecular weights of 47 kd, 44 kd, 38 kd, 28 kd, 20 kd and 15 kd, on a 15%-SDS polyacrylamide gel of the S-30 reactions with pRF2 or pRF4. To assign these protein products to their corresponding rib-specific ORFs, S-30 reactions were repeated using various available deletion derivatives, cat-insertion derivatives, and subcloned fragments of the 10 kb EcoRl cloned DNA (Figure 10). The results are shown in Table V.

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TABLE V

	RIB-SPECIFIC PROTEINS OBSERVED IN S-30 REACTIONS					
Plasmid		47,000	44,000	26,000	15,000	
- 1	Daltons (ORF)		Daltons (ORF	Daltons (ORF	Daltons (ÖRF	
-		3)	5)	4)	2)	
	pRF2	+	+	+	+	
-	pRF4	+	+	+	+	
-	pRF21	-	-	+	-	
	pRF5	-	•	•	+	
1	pRF29	-	-	-	•	
-	pRF12	+	-	+	+	
1	pRF10	-	-	-	-	
	pRF38	•	+	-	-	
	pRF24/pRF20	-	+	+	+	
Ì	pRF23	+	-	+	+	

Based on these results, protein products were assigned to ORF 3 (47 kd); ORF 5 (44 kd); ORF 4 (26 kd); and ORF 2 (15 kd), with molecular weights in close agreement with the predicted sizes.

The assignment of products to ORF 2 and the β riboflavin synthase gene were less straightforward than the assignments to the other ORFs. Since the S-30 reaction of pRF2 produced a 15 kd protein which was close to the predicted size of the proteins encoded by either gene, it was first assumed that this protein band actually contained both protein species. However, the cat insertion into ORF 2 in plasmid pRF38 completely removed this protein band, replacing it with a much smaller protein of 6 kd, which is in close agreement with the predicted size of the truncated ORF 2. Based on these results, the 15 kd protein appears to be generated only by ORF 2. It is not clear why the β riboflavin synthase protein is not visualized on the gels of the S-30 reactions. Taken in total, however, the results confirmed the existence of five rib -specific coding regions: ORF 5, ORF 4, ORF 3, ORF 2 and the β riboflavin synthase gene.

in addition, ORF 1 appeared to encode a 38 kd protein, while no product was identified for ORF 6.

Regulatory Mechanisms of the rib Operon

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In B. subtilis, a recurring pattern of gene organization and regulation for biosynthetic pathways has been observed by several investigators. The nucleotide sequences of the tryptophan biosynthetic pathway (Henner et al., Gene 34:169, 1984) and the de novo purine nucleotide pathway (Ebbole and Zalkin, J. Biol. Chem. 262:8274, 1987) of B. subtilis both contain clustered, overlapping genes transcribed as a polycistronic message and regulated at least in part by a novel transcription termination/anti-termination mechanism, involving a repressor protein which can be encoded by a gene unlinked to the biosynthetic operon (Zalkin and Ebbole, J. Biol. Chem. 263:1595, 1988). Since we found that the organization of the rib biosynthetic and regulatory genes is strikingly similar to those of the B. subtilis trp and pur pathways, we hypothesized that the rib operon might be regulated, at least in part, in a similar manner.

Briefly, the key characteristics of the transcription termination/anti-termination model include (Shimotsu et al., J. Bacteriol. 166:461, 1986): (i) the presence of a long 5' leader sequence that precedes the first gene in the operon; (ii) the presence in the RNA leader of two or more overlapping dyad symmetries that have the potential to form mutually exclusive RNA stem-loops, one structure functioning as a rho-independent transcription terminator and the other as an "anti-terminator" (blocking the formation of the rho-independent transcription terminator); (III) under repressive conditions, the repressor protein, activated by the end product of the pathway, binds to the nascent mRNA at a site which prevents formation of the anti-terminator, thus allowing formation of the terminator which terminates transcription; (iv) under derepressive conditions, binding of the unactivated repressor is precluded, resulting in the formation of the anti-terminator causing read-through transcription into the coding region of the operon.

As discussed above, the most likely site for initiation of transcription in the rib operon is a σ^A promoter, P₁, locat d about 290 bp upstream from the first given in the operon. Preliminary analysis of the RNA leader sequence indicated that it contained most, if not all, of the structures required for regulation by the

termination/anti-termination model. Within this region, a stem-loop structure followed by a string of thymidines resembling a rho-independent transcription terminator was identified approximately 50 bp upstream of ORF 5; this sequence has the potential to form a hairpin with a ΔG of -26 kcal/mol (Figure 9). In addition, several potential stem-loop structures with ΔG 's ranging from -13 to -16 kcal/mol were located within the rib 5 leader that could possibly qualify as the anti-terminator sequence.

In addition to the primary site for the initiation of transcription, usually located upstream from the first gene in the operon, there exist in some biosynthetic pathways secondary promoter sites located within the internal regions of the operon. The possibility of there being a second promoter site within the rib locus was also suggested by previous R-loop heteroduplex studies of the rib operon (Osina et al., FEBS Letters 196:75-78, 1986), showing two or more sites for the initiation of mRNA synthesis. Our preliminary analysis of the intercistronic gaps of the rib operon did not detect such secondary promoter sites. However, when this analysis was extended to all of the sequences within the operon, another σ^A promoter, P_2 , was identified within the 3 end of ORF 4, just downstream from the Sall restriction site (Figure 4). Thus it is possible that the expression of ORF 2, ORF 3, and the β -subunit for riboflavin synthase is also under the control of this secondary promoter. In addition, a possible third σ^A promoter, P_3 , was identified just upstream of ORF 2. Therefore ORF 2 is possibly also under the control of this additional promoter.

The location of putative coding regions, promoters and transcription termination sites on the DNA sequence of the 5.5 kb B. subtilis rib-specific region is shown in Table VI.

TABLE VI

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CODING REGIONS, PROMOTER, AND TRANSCRIPTION TERMINATION SITES OF THE B. SUBTILIS RIB OPERON				
		bp Number *		
Coding Regions	ORF 6 ORF 5 ORF 4 ORF 3 \$\beta\text{ riboflavinsynthase gene} \text{ ORF 2} ORF 1	364-678 1101-2183 2197-2841 2859-4052 4088-4549 4665-5036 5567-5057		
σ ^A Promoters	P ₁ P ₂ P ₃	771-799 2528-2556. 4545-4574		
rho -Independent Termination Sites	Upstream from 5' promoter Within 5' leader RNA At 3' end of rib operon	708-748 1034-1067 5038-5090		

a of Figure 3.

Example 8: Construction of vectors containing a modified rib operon

The above functional analysis of the rib operon of Bacillus subtilis for the first time delimiting the regulatory regions and open reading frames in the nucleotide sequence permits construction of new vectors which are useful for increasing the yield of riboflavin production. The knowledge of the location of the specific genes required for riboflavin biosynthesis, of the location of transcriptional control regions, and other relevant regions (e.g., RBS) in those genes allows changes in such regions to be made. There follow a few examples of such manipulations.

Construction of an Integration plasmid with a rib operon on a smaller DNA fragment

^b Coding region oriented in opposite direction.

The integrating vector used to construct the riboflavin overproducing strain RB50::[pRF8] contains a 10 kb EcoRl fragment including the rib operon. Since the rib operon appears to occupy less than 6 kb of DNA a new integration vector was constructed (pRF40) containing the rib operon on a smaller DNA fragment. The smaller size of this clone allows higher amplification of rib genes resulting in higher yields of riboflavin.

Referring to Fig. 12, pRF40 was constructed from pRF36 which is a plasmid in which the 0.8 kb Ncol fragment of pRF2 is replaced with a cat gene. The rib operon is contained on a 8.5 kb Xbal-EcoRl fragment. This fragment was isolated and ligated to pUC19 (Yanisch-Perron et al., Gene 33, 103, 1985; available from New England Biolabs, Boston, MA, USA and Bethesda Research Laboratories, Maryland, USA) digested with Xbal and EcoRl. The ligated DNA was transformed into DH5 α E. coli and plated onto LB plates containing 40 μ g/ml X-gal and 50 μ g/ml ampicillin. Analysis of miniprep DNA prepared from white colonies indicated that pRF39 contained the 6.5 kb Xbal-EcoRl fragment.

pRF39 was digested with EcoRI, treated with CIAP, and then ligated to a 1.6 kb EcoRI fragment containing the cat gene. The ligated DNA was then transformed into DH5 α E. coli and appropriate colonies selected for plating onto LB + 10 μ g/ml chloramphenicol; two colonies were chloramphenicol-resistant. Analysis of miniprep DNA prepared from these colonies confirmed the presence of the cat gene. One of these plasmids is pRF40 (Fig. 14).

Construction of plasmids containing transcriptionally modified rib operon

As described above, it is useful to replace the promoter and operator regions of the riboflavin operon with promoters allowing constitutive expression of the riboflavin biosynthetic genes. Plasmids containing such constructs can then be used to produce bacterial strains which will produce increased levels of riboflavin. A few examples, not limiting in the invention, are provided below.

Referring to Fig. 13, the riboflavin promoter and regulatory region were removed and replaced with an SPO1 promoter. We took advantage of the Bglll site located at position 1130 at the start of ORF3. Oligonucleotides were synthesized (RB5 and RB6, see Fig. 18) that recreated the DNA sequence 5 to the Bglll site (the first few amino acids of ORF5 and the SD sequence) up to position 1058. Reconstruction of the 5'-end of the operon stopped before any of the proposed DNA regulatory structures (Fig. 13). At their 5' ends the oligonucleotides contained BamHI, Nsil, and EcoRI restriction sites, allowing for placement of various promoters 5' to the rib operon. Because of the various restrictions sites in the rib operon it was necessary to construct the operon with the new promoters in several steps, as follows.

A 1.4 kb Sall-Bglll fragment was isolated from pRF36 (Fig. 13). This fragment was ligated with the two oligonucleotides and EcoRl-Sall-digested pUC19. The ligated mixture was then transformed into E. coli DH5 α cells and plated onto LB containing 50 μ g/ml ampicillin and 40 μ g/ml X-gal. Minipreps were prepared from Ap' white colonies; one plasmid having the desired structure is pRF46 (Fig. 13).

pRF46 was digested with BamHI and Sall and the 1.4 kb fragment isolated. This fragment was then ligated with the 400 bp EcoRi-BamHI fragment of pNH202 (pUC8 containing the SPO1-15 promoter, Lee and Pero, J. Mol. Biol., 152:247-265, 1981) and pUC19 cut with Sall and EcoRi. The ligated DNA was then transformed in DH5 α E. coli, which were plated onto LB + ampicillin + X-gal. Miniprep DNA was prepared from white colonies; and pPRF48 had the desired structure (Fig. 13).

pRF48 was digested with EcoRI and SaII and the 1.8 kb fragment isolated. This fragment was ligated with the 4.0 kb XbaI-SaII fragment (containing the rest of the rib operon) from pRF2 and XbaI, EcoRI-cut pUC19. The ligated mixture was then transformed into E. coli DH5α cells which were plated on LB + ampicillin + X-gal. Miniprep DNA was prepared from white colonles; pRF49 had the desired structure, and supernatants from culture containing this plasmid was yellow, indicating riboflavin production (Fig. 13).

To place the cat gene in pRF49, to allow selection in B. subtilis, the plasmid was digested with Xbal and ligated to a 1.3 kb cat-containing Xbal fragment from pEcc4. The ligated DNA was transformed in E. coli DH5 cells. Hundreds of Ap' colonies resulted, and the colonies were patched onto plates containing LB + 10 µg/ml chloromphenicol. Approximately 10% of the colonies grew on the chloramphenicol plates, indicating the presence of the cat gene. One cat-containing plasmid is called pRF50 (Fig. 14).

The above example shows placement of a new promoter upstream of ORFs. We found that it is also useful to place a promoter after P₂ between ORF3 and ORF4 in order to further increase riboflavin production. An example of such construction now follows.

Referring to Figs. 14 and 15, to place a copy of the SPO1-15 promoter upstream of ORF3 we made use of the restriction sites adjacent to the ORF4-ORF3 junction. The Clal site at position 2767 is located at the end of ORF4 and is unique in the rib operon. Another useful restriction site near the beginning of ORF3 is the Dral site at position 2892. Oligonucleotides were synthesized that recreat distriction the squence from the

above-mentioned Dral site past the start of ORF3 and placed a unique BaMHI site before the beginning of ORF3 (linkers P2-A and P2-B, Fig. 18). Another set of oligonucleotides recr ated the sequence from the Clal site past the end of ORF4 and placed an ECORI site at that location (linkers P2-CII and P2-DII, Fig. 18). The SPO1-15 promoter, located on a EcoRI-BamHI fragment, was then be placed between the BamHI and EcoRI sites created by the oligonucleotides. The entire operon was put together with this additional SPO1-15 promoter as follows.

Referring to Fig. 15, the 750 bp Sall-Bglll fragment containing the ORF4-ORF3 function was subcloned to pIC2OR (Marsh et al., Gene 32, 481-485, 1984). The resulting plasmid, pRF57, was then digested with Dral and Bglll, and the predicted 270 bp Dral-Bglll fragment was isolated. This fragment and linkers P2-A and P2-B were ligated to pIC2OR cut with Sall and Bglll. The linkers placed BamHl and Sall sites upstream of the 5 end of ORF3. (The sall site was chosen for convenience since Bglll and BamHl sites are compatible and will be removed later.) The ligation was transformed into E. coli DH5α cells. Plating onto LB medium + Amp and X-gal resulted in white colonies; pRF58 had the desired structure. The 330 bp Bglll-Sall fragment from pRF58 was isolated and ligated with 3.3 kb Bglll-Xbal fragment containing the 3 end of the rib operon from pRF36 (Fig. 12) and pUC19 cut with Xbal and Sall. The ligated DNA was then transformed into E. coli DH5α cells, resulting in white colonies; pRF62 (Fig. 15) had the desired structure. For convenience, the 3.6 kb BamHl-Xbal fragment was isolated from pRF62 and subcloned into BamHl-, Xbal-cut pUC19 (pRF64, Fig. 15). This plasmid now contained the 3.6 kb 3 end of the rib operon with an engineered BamHl site preceding ORF3.

To place the SPO1-15 promoter in front of the 3'-half of the rib operon containing the last three open reading frames, we digested pRF64 with EcoRI and BamHI and ligated it to a 400 bp EcoRI-BamHI fragment containing the SPO1-15 promoter. The ligated DNA was transformed into E. coli DH5 cells and miniprep DNA was prepared; pRF65 has the desired structure.

The SP01-15 promoter was than engineered to place a Clal site upstream of the promoter to reconstruct the end of ORF4. The EcoRl-BamHI fragment from pNH202 containing the SP01-15 promoter was ligated with linkers P2-ClI and P2-DII and pCl2OR-digested with BamHI and Clal. The ligated DNA was then transformed into E. coli DH5 α cells. White colonies resulted and miniprep analyses indicated that pRF63 had the desired structure. The 470 bp Clal-BamHI fragment was isolated then from pRF63 and ligated to the 2kb EcoRl-Clal fragment from pRF49 containing the SPOI-15 promoter and the 5 end of the rib operon and pRF64 (Fig. 15), containing the SPOI promoter and the 3 end of the operon, digested with EcoRl and BamHI. The ligated DNA was then transformed into E. coli DH5 α cells. Miniprep DNA was prepared; pRF66 had the desired structure. In addition, E. coli containing pRF66 produced small amounts of riboflavin on LB medium + ampicillin plates, confirming that the operon was still intact.

The last step was to ligate the cat gene into the unique Xbal sites of pRF66 as described above. The resulting plasmid, pRF69 (Fig. 15) contained the cat gene in the same direction as the rib operon.

To construct a plasmid containing the entire operon with the natural or wild-type ribP₁ promoter and the SPO1-15 promoter after ribP₂, the 6.3 kb EcoRI-BamHI fragment of pRF64, the 2.75 kb EcoRI-Clal fragment of pRF36, and the 470 bp Clal-BamHI fragment of pRF63 were ligated and tranformed into E. coli DH5 α cells. About 50% of the Apr colonies were yellow, indicating ribflavin production. Miniprep DNA was prepared from yellow colonies and pRF68 had the desired structure (Fig. 16). A cat gene was added to pRF68 at the Xbal site, as discussed above, to generate pRF71 (Fig. 16). This plasmid contained the cat gene in the same direction as the rib operon.

As another example of the construction of useful plasmids in this invention, there now follows an example in which one or more promoters can be introduced within the riboflavin operon without prior removal of existing DNA sequences.

As an example, a prototype modified operon was constructed in pRF78, which contains a single copy of the SPO1-15 promoter inserted within a 30 bp non-essential region located between ribP₁ and a putative rho-independent transcriptional termination site (Fig. 14), an inactivated ribP₁ promoter to prevent possible transcriptional interference of the SPO1-15 promoter, an active ribP₂ promoter, the five structural genes encoding rib biosynthetic enzymes, and approximately 1.5 kb of flanking DNA nucleotide sequences downstream from the end of the riboflavin operon.

Referring to Fig. 14, the 1.7 kb Ncol-Psti fragment of pRF2, a fragment that contains the 5' promoter region of the rib operon and flanking regions, was first subcloned into mp19, a derivative of the E. coli bacteriophage vector M13 (United States Biochemical Catalog, 60-61, 1987; available from New England Biolabs, Massachusetts, USA). One recombinant phage, M1.7, was recovered and standard DNA sequence analysis of the promoter region revealed a spontaneous mutation of the -10 region of the ribP₁ promoter, a TA-to-CT change, which may inactivate the promoter. Single stranded DNA was prepared and annealed to a synthetically-generated 55 bp DNA oligomer (see Fig. 17), containing a combination of restriction enzymes

sites, 5'-EcoRi-Smal-BamHi-3', flank d on either side by additional sequences homologous to the DNA region upstream from ribP₁. Double-stranded DNA molecules were synthesized using standard site-dir ct d mutagenesis (SDM) protocols. These DNA molecules were introduced into the E. coli host TG-1 (Amersham Corp., Illinois, USA) by transfection to generate recombinant phage plaques. One recombinant phage was found to contain the desired modified DNA sequence, as determined by standard DNA sequence analysis.

The modified rib promoter region was then rejoined to the rib structural genes of the operon using a pair of unique Nsil restriction enzymes sites 750 bp apart of the flank the ribP₁ region and surrounding sequences. Double-stranded DNA molecules of the phage recombinant were prepared, digested with Nsil, the 750 bp fragment isolated, and the fragment ligated to dephosphorylated, 8.7 kb Nsil fragment of pRF39ΔR1 (a plasmid derived from pRF39. Fig. 12, that contains the wild-type rib operon). The ligated DNA molecules were introduced into E. coli DH5α cells by transformation, selecting for ampicillin-resistance, which resulted in the recovery of an Ap' colony harboring the desired recombinant plasmid, pRF75.

The SPO1-15 promoter was next inserted upstream from ribP₁ by digesting pRF75 with a combination of EcoRl and BamHl enzymes, ligating the cut DNA to purified 400 bp EcoRl-BamHl SPO1-15-containing restriction fragment, and introducing the ligated DNA into E. coli DH5α cells by transformation, selecting for ampicillin-resistance. One Ap^r colony was found to harbor the recombinant plasmid, pRF77, containing the desired SPO1-15-modified rib operon. A chloramphenicol-resistance gene, cat, on a 1.6 kb Xbal restriction fragment, was subsequently introduced into pRF77 at the unique Xbal site, generating plasmid pRF78 (Fig. 14).

This prototype operon was further modified to contain an active ribP₁ promoter, and/or a second copy of the SPO1-15 promoter introduced downstream from ribP₂ within a intercistronic region between the rib coding regions ORF3 and ORF4, as described above. For example, plasmid pRF88, containing a derivative of the modified rib operon in pRF78 with an active ribP₁ promoter (Fig. 14) was constructed by the same procedure described above, using a recombinant phage containing the wild-type ribP₁ promoter. In other examples, a second copy of the SPO1-15 promoter, located downstream from ribP₂, was inserted into the existing modified rib operon-containing plasmids pRF78 and pRF88 by removing the 2.0 kb Bglll fragment of either plasmid DNA and inserting the 2.4 kb Bglll fragment of pRF66, generating plasmids pRF81 and pRF89 respectively (Fig. 14).

Construction of Ade RB50 strains

It is important to use strains of bacteria that require as few components to be added to a fermentation medium as possible. Such strains are cheaper to ferment in order to produce riboflavin. To this end, adenine revertants which contained amplified modified rib operons were constructed. These revertants may not be true revertants of pur-60, but rather include mutations at another site which suppresses the requirement for adenine. As discussed below they produce about 25% more riboflavin than the non-reverted strains. Examples of such constructions are now described.

Plasmids pRF8, pRF40, pRF50, pRF69, pRF71, pRF78, pRF81, pRF88 and pRF89 were each transformed into RB50 (a RoF¹, deregulated B. subtilis strain) selecting for chloramphonicol resistance (Cm²). A resistant colony was chosen for each strain. Ade revertants of each strain was isolated by growing bacteria in RMM1 broth containing 10 μg/ml adenosine, and plating samples of the cultures onto minimal agar plates. One colony from each Ade strain was selected and the vector DNA was amplified by selecting colonies that grow on increasingly higher levels of chloramphenicol, to a maximum level of 60 μg/ml.

Second site Integration

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As described above, it is important to amplify an engineered rib operon in the B. subtilis chromosome to achieve high titers of riboflavin. It is also important to ensure that the number of DNA copies of the rib operon within a chromosome are not limiting to riboflavin production. Further amplification of the rib operon can be achieved by integrating and amplifying copies of the rib operon at more than one site in the B. subtilis chromosome to further increase riboflavin yield. One example of how such second site integration can be achieved is described below.

The above described vectors have all relied upon the cat gene to allow integration at the site of the rib operon. In order to insert the rib genes at a second site, it is pr f rable to have a differ nt antibiotic resistance gen for us at that second site. For example, a tetracycline-resistance (tet) from B. subtilis can be used (Perkins and Youngman, J. Bacteriol., 155:607-615, 1983). Such tet genes are will known to those

of ordinary skill in the art and are readily available to such persons. In one such construction, for example, pRF78 (Fig. 14), which contains a modified version of the rib operon, the plasmid can be cut with Xbal and ligated to a 2.4 Xbal fragment containing the tet gene. The resulting plasmid contains the tet gene at the Xbal site and is called pRF85.

A strain which is deleted for the entire rib operon and which has a tet gene integrated at a second site is required to cause integration of pRF85 at that site. One such site is the bpr gene encoding bacillopeptidase F, a minor non-essentiall extracellular protease. An E. coll plasmid containing the bpr gene, pKT2, (Sloma et al., J. Bacteriol., 172:1470-1477, 1990) was digested with EcoRV. This EcoRV site is in the coding region of bpr. The DNA was then ligated to a 2.4 kb Eco RI fragment containing the tet gene that had been blunt-ended. The resulting plasmid (containing the tet gene at the EcoRV site of bpr) was called pKT2-tet. This DNA was linearized with EcoRI and then transformed into RB52, a strain deregulated for riboflavin synthesis. Tet colonies resulted and one such colony was called RB54. The integrated tet gene at bpr will function as a homologous sequence for the Integration of pRF85.

To ensure that the cloned riboflavin operon of pRF85 would be inserted at the second chromosomal site containing the tetracycline-resistance gene, a region containing the original riboflavin operon and flanking DNA, equalling that contained in pRF85, was deleted from the chromosome of RB54 by in vitro methods. Briefly, this involves first generating an E. coli recombinant plasmid where the cloned riboflavin operon and flanking regions between the Ncol and Xbal restriction sites are removed and replaced by a chloramphenicol-resistance gene, cat, that is expressed in B. subtilis bacteria. This plasmid is then used to delete the chromosomal riboflavin operon by transforming RB54 with linearized plasmid molecules and selecting for chloramphenicol resistant (Cm^r) bacteria. Cm^r bacteria result from a recombinant event (marker-replacment) which replaces the wild-type rib genes with the deleted copy containing the cat gene.

Specifically, plasmid pRF34 (see example 6) was used to generate an E. coli plasmid containing an in vitro-generated riboflavin operon deletion. This plasmid is derived from pRF2 where the riboflavin operon is flanked on either end by two unique Xbal sites (one site located upstream from the 5'-end of the rib operon next to the deleted 0.8 kb Ncol fragment and the second site located approximately 1.6 kb downstream from the end of the operon) and a cat gene is inserted outside of this region. By digesting pRF34 with Xbal and ligating the cut DNA molecules under dilute DNA concentrations, a recombinant plasmid, pRF82, was recovered where a 7.2 kb region containing the riboflavin operon is removed and essentially replaced with the cat gene. Plasmid pRF82 was linearized by restriction enzyme digestion and the cut DNA used to remove the chromosomal riboflavin operon of RB54 by DNA transformation, selecting for Cm' bacteria, resulting in marker replacement. Cm' colonies were screened for riboflavin auxotrophy and one Rib-Cm' colony, RB55, was recovered for further investigation.

Plasmid pRF85 was transformed into strain RB55, selecting for Rib*. One Rib* transformant was chosen and called RB58. This strain has the rib operon integrated at bpr by homologous recombination between the tet' genes in the plasmid and the chromosome. Chromosomal DNA from RB58 was prepared and can be used to transform RB50::[pRF69], selecting for Tet'. These resistant colonies will then have the modified rib operon integrated at the site of the rib operon and at bpr. The rib operon integrated at rib will be amplified by selecting for colonies that grow in the presence of increasing levels of chloramphenicol as described above, and the second copy of the rib operon will be amplified by selecting colonies that grow on increasing levels of tetracycline.

Example 9: Fermentative Production of Riboflavin

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Evaluation of riboflavin-overproducing strains was conducted in Chemap 14-liter vessels in carbon-limited fed-batch fermentations, with riboflavin content measured by HPLC. Since enzymes encoded by the genes for riboflavin synthesis are rate-limiting, the rib genes, which were amplified, were maintained at high-copy number by the inclusion of $60~\mu g/ml$ chloramphenicol in the inoculum seed train, but not in the fermentary

A culture of B. subtilis RB50::[pRF69] was grown on Tryptose Blood Agar Base (TBAB Difco) containing 60 μg/ml of chloramphenicol (CAM). Colonies were transferred to 300 ml baffled flasks containing 25 ml of riboflavin minimal medium (RMM; containing sodium glutamate 2.0 g/l, Casamino acids (Difco) 0.2 g/l, Yeast extract (Difco) 0.2 g/l, KH₂PO₄ 6.0 g/l, K₂HPO₄ 14.0 g/l, (NH₄) ₂SO₄ 2.0 g/l, sodium citrate 1.0 g/l, MgSO₄•7H₂O 0.2 g/l, adenosine 0.05 g/l, glucose 15.0 g/l, pH 7.0) with 60 μg/ml CAM. The inoculated flasks were incubated by shaking at 250 rpm and 37 °C. After 8 hours, sterile glycerol was added to a final concentration of 15% and 1 ml aliquots w re stored at -80 °C.

In order to initiate a ferm ntation a frozen vial of RB50::[pRF69] was thawed at 37°C and transferred

into a 300 ml baffled flask with 25 ml of RMM with 60 µg/ml CAM and shaken at 250 rpm and 37 °C. After 8 hours, 6 ml of the growing culture was used to inoculate 300 ml of fermentation medium (see Table VII below) in a 2 liter transfer flask. Such flask contained 300 ml of fermentation medium to which had been added 90 ml of a mixture of 15% glucose and 30% maltose. Chloramphemicol was added to a final concentration of 60 µg/ml. After incubation for 12 hours at 200 rpm on a shaker with a 2" diameter orbit at 37°C, the contents of such a flask was transferred to 7 liters of fermentation medium in a 14 liter fermentation vessel.

During fermentation, the broth was continually monitored for pH and dissolved oxygen (DO₂). Off gas was continuously analyzed by quadrapole mass spectrometry and carbon dioxide evolution (CER) and oxygen uptake rates were recorded.

A comparison of several fermentations demonstrated the reproducibility of the control systems. The initial carbohydrate was exhausted from fermentation with RB50::[pRF8]₅₀ after 4 hours of growth, causing a rise in pH and a fall in CER. At that point, carbohydrate feeding was initiated and logarithmic growth resumed until DO₂ became limiting at 6 hours. The rate of carbohydrate feeding was computer-controlled to maintain the DO₂ between 10-20% of saturation throughout the remaining fermentation time.

Excess carbohydrate in the fermentors does lead to oxygen starvation and reduced riboflavin production. Oxygen transfer limitations determine the duration of logarithmic growth, final cell density and the riboflavin production rate. To increase the oxygen transfer rate, Chemap fermentors were run at 1000 rpm with a head pressure of 0.6 atmospheres.

Supplementation of the medium carbohydrate feed with yeast extract led to an increase in riboflavin production as compared to media without supplementation (Figure 11, open squares: RBF-14; Table VII). However, because of its high cost, the amount of yeast extract was systematically reduced by substituting less expensive, inorganic ingredients. Substitution of ammonium hydroxide for sodium hydroxide in pH control allowed a reduction of yeast extract in the feed and resulted in an increase in both cell mass and riboflavin titer (Figure 11, closed squares: RBF-22; Table VII). Fermentation times were also reduced. In other fermentations, moreover, yeast extract was completely eliminated from the feed and replaced with a combination of inorganic salts of ammonium and phosphate, resulting in a further increase in riboflavin production and a reduction of process time (Figure 11, open circles: RBF-23; Table VII).

The original RB50::[pRF8]₅₀ was auxotrophic for adenine because of its pur-60 mutation. When experiments were conducted to determine the minimum amount of adenosine required by the strain, in order to minimize its inhibition of earlier biosynthetic enzymes involved in the pathway leading to the ribofiavin-precursor IMP (Figure 2), RB50::[pRF8]₅₀ (and, In general, RB50 strains with a rib operon amplified within their chromosome) was found to be unstable in its adenosine requirement and prototrophic revertants (Ade^{*}) were produced at a fairly high frequency. In shake flasks, the Ade^{*} revertants appeared to grow and produce riboflavin at least as well as the RB50::[pRF8]₅₀ parent. When evaluated in fermentors, the revertant, RB50::[pRF8]₅₀(Ade^{*}), did not require adenosine in the media formulation. More importantly, the revertant grew-at a faster rate and produced 25% more riboflavin than its parent strain in less time. A titer of 5.4 g/l riboflavin was produced in 49 hours (Figure 11, closed circles: RBF-29; Table VII). In additional fermentations, moreover, Hy Soy T was removed from the initial charge or medium and replaced with corn steep liquor resulting in a further increase in riboflavin production to 6.3 g/l in 48 hours. (RBF-42, Table VII).

Under these fermentation conditions, further significant increases in riboflavin production were demonstrated using bacterial strains that contained engineered riboflavin operon DNA. Strains containing the wild-type riboflavin operon on a 6.5 kb EcoRl-Xbal restriction fragment, RB50::[pRF40]₆₀(Ade^{*}), produced 7.4 g/l of riboflavin in 48 hours. Moreover, strains containing a transcriptionally-modified rib operon where the rlbP₁ promoter and regulatory region were replaced by the constitutive SPO1-15 promoter, RB50::[pRF50]₆₀(Ade^{*}), produced 9.0 g/l of riboflavin in 48 hours. These results demonstrate that modification of the riboflavin operon through the removal of regulatory regions and/or through the introduction of stronger, constitutive exogenous promoters leads to increases in riboflavin titer.

TABLE VII

	FERMENTATION COMPONENTS AND CONDITIONS					
5	Component	RBF-14	RBF-22	RBF-23	RBF-29	RBF-42
	Initial Charge (g/l)					
	Glucose	10.00	15.00	15.00	15.00	15.00
	Corn step liquor	· -	_	-	-	10.00
10	Hy Soy T	15.00	15.00	15.00	10.00	
	Sodium glutamate				5.00	5.00
	Amberex 500	15.00	15.00	20.00	20.00	20.00
	KH₂PO₄	5.00	5.00	7.50	7.50	7.50
	MgGl ₂ .6H ₂ O	0.5	0.5	1.50	1.50	1.50
15	MnSO ₄	0.05	0.05	0.05	0.05	0.05
	Adenosine	0.05	0.05	0.05	-	· -
	MAZU DF37C	2.50	2.50	2.50	2.50	2.50
	FeCl ₃	-	_	0.025	0.02	0.02
	CaCl ₂		<i>≟</i>	0.50	0.50	0.50
20	ZnSO ₄		-	0.0005	-] -
	CuCl ₂	-	-	0.0013		-
	CoCl ₂	-	-	0.0013	-	-
	Nutrient Feed (g/l)					
25	Amberex 500	160.00	120.00			
	NH₄CI		-	7.50	7.50	7.50
	(NH ₄) ₂ SO ₄	-	_	7.50	7.50	7.50
	KH₂PO₄		-	15.00	15.00	15.50
	MgSO₄.7H₂O	-		2.50	2.50	2.50
30	DL-70 syrup (as DS)	600.00	600.00	600.00	600.00	600.00
	pH Control Range					
	6.6	H₂SO4	H₂SO4	H₂SO ₄	H₂SO₄	H₂SO₄
	6.5	NaOH	NH₄OH	NH4OH	NH₄OH	NH ₃
35	Conditions	1				
	Air (vvm)	1.0	1.5	1.5-2.0	1.5	1.50
	RPM	1000.0	1000.0	1000.0	1000.0	1000.0
	Temp. (*C)	37.0	37.0	37.0	37.0	37.0
40	Pressure (bar)	0.5	0.5	0.5-0.75	0.6	0.6
	Riboflavin (g/l)	3.4	4.1	4.3	5.4	6.3
		(64 hrs)	(56 hrs)	(53 hrs)	(49 hrs)	(48 hrs.)
	Dry Weight (g/l)	33.6	· 36.0	36.8	ND	44.6

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The kinetics of riboflavin production in the various fermentations were analyzed using the Luedeking-Piret model. In all cases, the specific productivity declined from the conclusion of the exponential growth phase to the end of fermentation. Also, it was clear that riboflavin production was growth-associated under the fermentation conditions used.

We have discovered that the yield of riboflavin can be increased by changing the fermentation components and conditions. The yield of riboflavin can be increased compared to those conditions described above using those fermentation components and conditions shown in Table VIII.

TABLE VIII

	(g/liter)	
Initial Batch		
Yeast Extract	20	
Glucose	25	
KH₂PO₄	7.5	
MgCl ₂ .H ₂ O	1.5	
CaCl ₂ .2H ₂ O	1.0	
MnSO ₄	0.05	
FeCl ₃ .6H ₂ O	0.025	
Mazu DF37C	2.5	
Corn Steep Liquor	10	
Sodium Glutamate	5	
Feed Medium (3 liters total used)		
Glucose	583.3	
NaCitrate	6.67	
KH₂PO₄	15	
Succinic Acid	1.67	
MgSO₄.7H₂O	1.67	

25

10

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Briefly, in one such fermentation the starting material is 6.65 liters of batch medium and 0.35 liters of bacterial (RBSO::[pRF50]₅₀Ade) inoculant. Oxygen levels are monitored with a Chemap polarographic dissolved oxygen electrode. Dissolved oxygen levels are maintained at 15% ± 5% by means of computer regulated addition of the feed medium. Total feed added is about 3.0 liters in 48-56 hours. Fermentation pH is maintained at 6.5± 0.1 (using 1N H₂SO₄ and NH₃ gas), and fermenter pressure is maintained at 0.6 bars, and air flow at 10.5 liters/min. Under these conditions, strain RB50::[pRF50]₆₀(Ade) produced 11.0 g/l riboflavin in 48 hours, which represents an improvement in production of approximately 20% compared to the previous fermentation conditions. Finally, a further increase in riboflavin production was demonstrated using the bacterial strains RB50::[pRF69]₆₀(Ade) containing a transcriptionally-modified riboflavin operon containing two SPO1-15 promoters, one replacing ribP₁ and regulatory sequences, and a second inserted between ORF3 and ORF4. This strain produced 13.0-14.0 g/l riboflavin in 48 hours, and 15 g/l in 56 hours, demonstrating that increased transcription of the riboflavin operon using two strong exogeneous promoters increases production levels of riboflavin.

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Deposit of Microorganism

Plasmid pRF69 has been deposited with the American Type Culture Collection on June 6, 1990 under the Budapest Treaty and was assigned Accession No. ATCC 68338.

E. coli strain DH5 containing plasmid pRF50 has been deposited with the American Type Culture collection on May 30, 1990 under the Budapest Treaty and was assigned Accession No. ATCC 68332.

E. coli strain DH5 alpha containing plasmid pRF78 has been deposited with the American Type Culture collection on May 30, 1990 under the Budapest Treaty and was assigned Accession No. ATCC 68333 and Bacillus subtilis strain RB58 has been also deposited with the American Type Culture collection on May 30, 1990 under the Budapest Treaty and was assigned Accession No. ATCC 55053.

Bacillus subtilis strain RB50 was deposited on May 23, 1989 with the Agricultural Research Culture Collection (NRRL), Peoria, Illinois, under the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedures, and was assigned accession number B 18502.

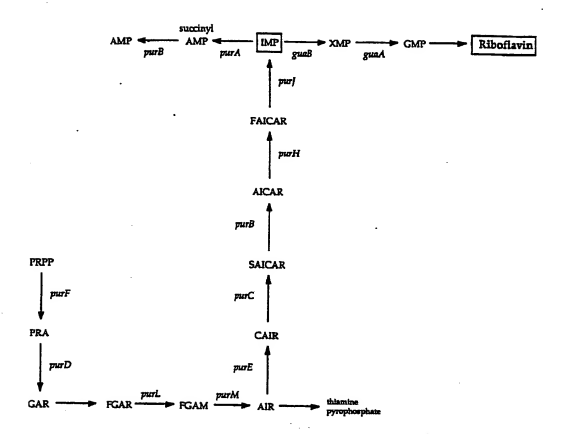
Claims

- 1. A v ctor comprising first a nucl ic acid sequence of bacterial or y ast origin coding for one or more riboflavin biosynthetic proteins and second one or more transcription elements not naturally associated with this nucleic acid sequence.
- 5 2. A vector according to claim 1 wherein said nucleic acid sequence of bacterial origin is derived from Bacillus especially Bacillus subtilis or E. coli.
 - 3. A vector according to claim 1 or 2 wherein said nucleic acid sequence is coding for at least 5 riboflavin biosynthetic proteins.
 - 4. A vector according to any one of claims 1 to 3 which comprises one or two such transcription elements.
- 5. A vector according to any one of claims 1 to 4 wherein said transcription element is a promotor.
 6. A vector according to claim 5 wherein said promotor is a promoter chosen from a promoter naturally associated with a gene from SPO1 bacteriophage, a promoter naturally associated with veg, army; and apr, and other transcription regulatory elements activated by the gen product of sacQ.
 - 7. A vector according to claim 6 which is pRF50, pRF69, pRF70, pRF71, pRF78, pRF81 or pRF89.
- 8. A recombinant bacterium comprising at least one copy of an exogenously introduced nucleic acid sequence at one or more sites within its chromosome, which sequence encodes one or more riboflavin biosynthetic proteins and is heritable and capable of expression by the bacterium, such that riboflavin biosynthesis by the bacterium is increased relative to a bacterium lacking such nucleic acid sequence.
- 9. A recombinant bacterium comprising a bacterium which has been transformed by a vector according to any one of claims 1-7 whereby at least one copy of said nucleic acid sequence including said transcription elements has been introduced at one or more sites within its chromosome and said nucleic acid sequence including said transcription elements is heritable and capable of expression by the bacterium, such that riboflavin biosynthesis by the bacterium is increased relative to a bacterium lacking such nucleic acid sequence including said transcription elements.
- 10. A recombinant bacterium according to claim 9 wherein said nucleic acid sequence including said transcription elements is present in a plurality of copies at such sites.
 - 11. A recombinant bacterium according to claim 9 or 10 wherein said nucleic acid sequence including said transcription elements has been introduced at one or two sites within its chromosome.
 - 12. A recombinant bacterium according to any one of claims 8 to 11 wherein said bacterium is deregulated for riboflavin gene expression.
 - 13. A recombinant bacterium according to any one of claims 8 to 12 wherein said bacterium is an E. coli or Bacillus, especially Bacillus subtilis strain.
 - 14. A recombinant bacterium according to claim 13 wherein said B. subtilis strain is RB50 or RB58.
- 15. A process for the production of riboflavin characterized by growing a recombinant bacterium according to any one of claims 8 to 14 under suitable growth conditions.
 - 16. A process according to claim 15 wherein such suitable growth conditions are characterized by limiting the availability of a component of the growth medium in such a way that aerobic conditions for the growth of said recombinant bacterium are maintained.
 - 17. A process according to claim 16, wherein said component is chosen from a carbon source, nitrogen source or a component required by said recombinant bacterium.
 - 18. A process according to claim 17, wherein said component is glucose, or a carbonic acid.
 - 19. A process according to claim 16, wherein said limiting step comprises limiting introduction of said component in a feed medium.
 - 20. A process according to claim 19, wherein said component is glucose, or a carbonic acid.

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FIGURE 2.



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FIGURE 3A

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FIGURE 3B

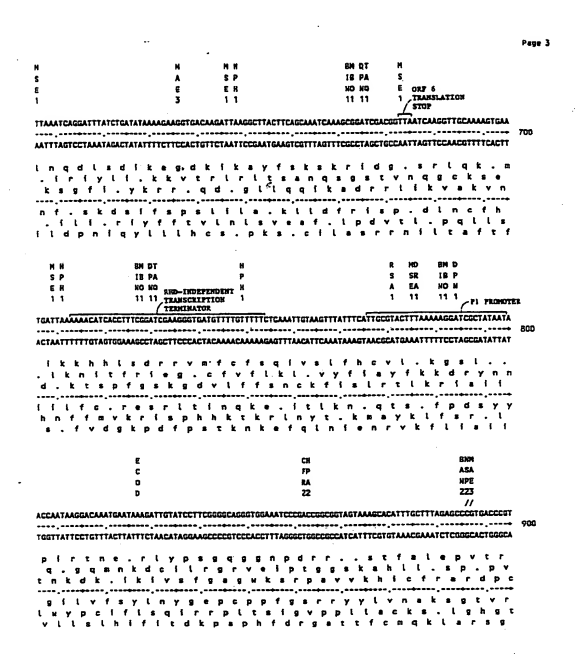


FIGURE 3C

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FIGURE 3D

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FIGURE 3E

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FIGURE 3F

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ilyrrophkrroecvififfpk.seshvykckf
dflasmsk.spyrlrpfsfvaklkrisci.vq
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FIGURE 3G

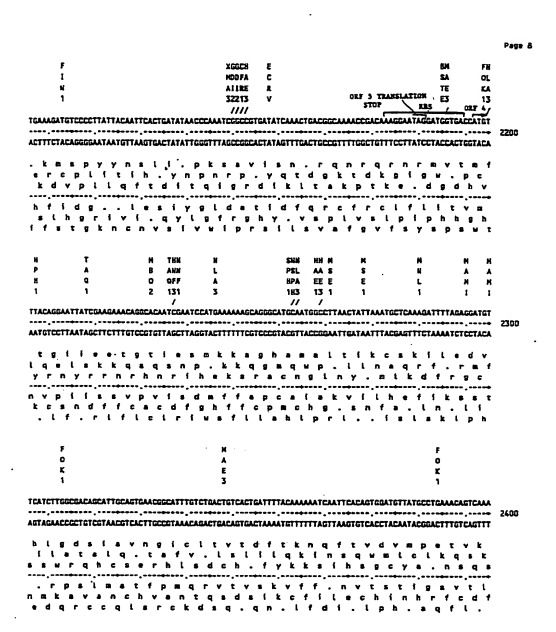


FIGURE 3H

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GACGGAACTGCGGA G g t a e t e t r k - r n c g	TAGTGCTTAACTICTC	P2 PROMOTE MAAGCAACGCAGTTTAG TITTTGGTTGCGTCAAATG k s n a v y k a t q f t k q r's l l f l l a t .	EXECUTATION TO THE PROPERTY OF	m d p :	AGTAATTGTTT sltk h.qk ink:	TTGTAACCAAAATTI tlvlk hwf.i nigfk	2600
CTGCCTTGACGCCT d g t a e t e t r k r n c g	TAGTGGGTTAAGTGTGTGTGTGTGTGTGTGTGTGTTAAGTGGTTAAGTTGTG	P2 PROMOTE AMAGCAGGAGTTTAG TITTEGTTGCGTCAAATG k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v	EXECUTION TO THE PROPERTY OF T	m d p :	AGTAATTGTTT sltk h.qk ink:	TTGTACCAAAATTI tlvlk . h w f . i n i g f k	2600
GACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e t r k r n c g	TAGTGCTTAACTICTC	P2 PROMOTE AMAGCAGGAGTTTAG TITTEGTTGCGTCAAATG k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v	EXECUTION TO THE PROPERTY OF T	m d p :	AGTAATTGTTT sltk h.qk ink:	TTGTACCAAAATTI tlvlk . h w f . i n i g f k	2600
GACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e l r k r n c g s p v e s v s s r	TAGTGCTTAACTICTC itriee cshelkr nhtn.rs iivriss fdesnfl	P2 PROMOTE MANAGEMEGRATITAGE K S n a V y k a t q f t c k q r s l l f l l a t . f a V c n v s f c r l k	ERATGATTTAAATTI y d i k m i . i i . f k . s k i y i i . s h n i	TTACCTGGCC m d p : c w t r n g p v ' { g g f h v r	AGTAATTGTTT s l t k h q k i n k d n v f	TIGTACCAAAATTI tivik hwf.i nigfk vntki fcqn. fmpki	2600
GACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e t r k r n c g \$ p y e s r f q p	TAGTGCTTAACTICTC itriee csheikr nhtn.rs iivriss fdcsnfi f.vfql	P2 PROMOTE MANAGEAGGEAGTTTAG ITTTEGTTGCGTCAAATC k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v s f c r l k	EXECUTION TO THE PROPERTY OF T	MATEGACCECT TTACCTEGGC m d p : c w t r n g p v f { E g f h v r f p g t	AGTAATTGTTT s l t k h . q k i n k i n v f c t m l l	TTGTACCAAAATTI tivik .hwf.i nigfk vntki fcqn. fmpkl	2600
GACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e t r k r n c g	TAGTGCTTAACTICTC itriee cshelkr nhtn.rs iivriss fdesnfl	P2 PROMOTE MANAGEMEGRATITAGE K S n a V y k a t q f t c k q r s l l f l l a t . f a V c n v s f c r l k	ERATGATTTAAATTI y d i k m i . i i . f k . s k i y i i . s h n i	MATEGACCECT TTACCTEGGC m d p : c w t r n g p v f l g g f h v r f p g t	AGTAATTGTTT s l t k h q k i n k d n v f	TTGTACCAAAATTI tivik .hwf.i nigfk vntki fcqn. fmpkl	2600
GACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e t r k r n c g	TAGTGCTTAACTICTC itriee csheikr nhtn.rs iivriss fdcsnfi f.vfql	P2 PROMOTE MANAGEAGGEAGTTTAG ITTTEGTTGCGTCAAATC k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v s f c r l k	ER ETATGATTTAAA TATACTAAATTI y d i k m i . i i . f k . s k / f i . s h n l	MATEGACCECT m d p : c w t r n g p v f i E g f h v r f p g i	AGTAATTGTTT s l t k h . q k i n k i n v f c t m l l	TTGTACCAAAATTI tlvlk h w f . i n i g f k V n t k i f c q n . f m p k l	2600
CACCGCACTGACGCCI d g t a e t e l r k r n c g s p v s r r f q p	TTAGTGTGCTTAAGTCTC itriee csheikr nhtn.rs fitries fdcsnfi f.vfqi AfM LOS	P2 PROMOTE AMAGCAAGGAGTTAG ITTTTGGTTGGGTCAAATC k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v s f c r l k HH -8 NA 8	ER ETATGATTTAA TATACTAAATTI y d i k m i i i f k . s k , f i x h n i H ION A BB	MATGGACCCC m d p: c w t r n g p v li g g f h v r f p g t D H P S N E	AGTAATTGTTT s l t k h . q k i n k: d n v fc t m l l	TTGTAACCAAAATTI t (v k . h w f . i n i g f k v n t k i f c q n . f m p k t D ND P BP R ON	2600
CTGCCTTGACGCCT d g t a e t e l r k r n c g S p v s r r f q p	TAGTGCTTAACTCTC itriee csheikr htn.re itvriss fdcsnfi f.vfqi AFM LOS UKE	P2 PROMOTE AAAAGCAACGCAGTTTAG ITTTTGGTTGGGTCAAATG k s n a v y k a t q f t k q r's l l f k q r's l f c r l k HH -8 NA 8 FE V	EXECUTATION TO THE PROPERTY OF	MATGGACCCC m d p: c w t r n g p v li g g f h v r f p g t D H P S N E	AGTAATTGTTT s l t k h . q k i n k i n v c t m l l	TTGTAACCAAAATTI t (v k . h w f . i n i g f k v n t k i f c q n . f m p k t D ND P BP R ON	2600
GACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e t r k r n c g s p v e s v s s r r r f q p p P R	TAGTGGGTTAACTICTC itriee cshelkr htn.rs ivriss fdcsnfl f.vfql AFM LOS UKE 111	P2 PROMOTE AMAGCAACGCAGTTTAG K S n a V y k a t q f t k q r's l l f l l a t . f a V c n v s f c r l k HH	ER ETATGATTTAAATTI y d i k B i . i i . f k . s k i y f i . s h n i H MM A BB E CO 3 21	MATEGACCECT m d p : c w t r n g p v f i E g f h v r f p g t D M P S N E 1 1	AGTAATTGTTT s l t k h . q k i n k i n k t m l l	TTGTAACCAAAATTI tivik h w f . i n i g f k v n t k i f c q n f m p k l D ND P B P N ON 1 11	- 2600 F
GACGGAACTGCGGA d g t a e t e t r k r n c g s p v a s v s s r r r f q p	TAGTGCGTGAGGGCTTAACTACCA	P2 PROMOTE AMAGCAAGGAGTTAG k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v s f c r l k HH 8 NA 8 FE V 33 2 TATTCGGCCTGACAGAAG	ER ETATGATTTAAA PATACTAAATTI y d l k m i l l f k . s k / f i l s h n l H HOR A BB E CO 3 21	TTACCTEGGC	AGTAATTGTTT s l t k h . q k i n k i n v f c t m l l B i OI	TTGTAACCAAAATTI t l v l k h w f . i n i g f k v n t k i f c q n . f m p k l D	2600
CACGGAACTGCGGA d g t a e t e t r k r n c g s p v e s t v s e r r f q p	TAGTGGCGTGAGCTTAACTA A FM L OS U KE 1 11	P2 PROMOTE AAAAGCAACGCAGTTTAG TITTTCGTTGCGTCAAATC k s n a v y k a t q f t k q r's l l t k q r's l t t q r's l t t q r's l t k q r r r r r r r r r r r r r r r r r r	ERTAGATTAAATTI y d i k	TTTACCTGGGC	AGTAATIGTTT s l t k h q k i n k i n k i n t l d n v f t m l l	TTGTAACCAAAATTI t l v l k . h w f . i n i g f k V n t k f c q n . f m p k l D H D P B P R O H 1 1 1	- 2600 r
CACGGAACTGCGGA CTGCCTTGACGCCT d g t a e t e l r k r n c g S p v e s d v s e r r f q p	TAGTGCGTGAGGGCTTAACTACCA	P2 PROMOTE AAAAGCAACGCAGTTTAG TITTTCGTTGCGTCAAATC k s n a v y k a t q f t k q r's l l t k q r's l t t q r's l t t q r's l t k q r r r r r r r r r r r r r r r r r r	ERTAGATTAAATTI y d i k	TTTACCTGGGC	AGTAATIGTTT s l t k h q k i n k i n k i n t l d n v f t m l l	TTGTAACCAAAATTI t l v l k . h w f . i n i g f k V n t k f c q n . f m p k l D H D P B P R O H 1 1 1	- 2600 r
d g t a e t e i r k r r n c g s p v e s d v s s r r r f q p GATCAATTACTGTG	TTAGTGTGCTTAACTTCTC itrfee cshelkr nhtn.rs ivriss fdesnfl f.vfql AFM LOS UKE 111 / GATGGCGTGAGCTTAACCA	P2 PROMOTE AMAAGCAAGGAAGTTAA k s n a v y k a t q f t k q r s l l f l l a t . f a v c n v s f c r l k HH	ERATGATTTAAATTI y d i k	TTACCTGGCC m d p : c w t r n g p v ? (R g f h v r f p g : D H P S H E 1 1	AGCAGGATAGG	TTGTAACCAAAATTI t v k h w f . I n i g f k v n t k f c q n f m p k D ND P B P N O N 1 1 1 ATCAGCGAAACCATC	- 2600 r
CEACGGAACTGCGGA d g t a e t e t r k r r n c g sp y e s d y s s r r r f q p GATCAATTACTGTG CETAGTTAATGACAC	TAGTGGCGTGAGCTTAACTA A FM L OS U KE 1 11	P2 PROMOTE AAAAGCAAGGAAGGAAGAAGAAGAAGAAGAAGAAGAAGAA	EX ETATGATTTAA LATACTAAATTI y d t k m f . 1 t f k . s k 1 y f i i s h n t A BB E 00 3 21 ACACAGTEACT	TTACCTEGGC	AGTACTATTOTT s l t k h q k i n k i n k i n c t m l l MI B i O TACCGCATACG ATGGCGTATGC	TTGTAACCAAAATTI t l v l k h w f . i n i g f k v n t k i f c q n . f m p k l D H D P B P N O N 1 11 ATCAGCGAAACCATC	- 2600 r
CTAGGTAATTACTGTG	TTAGTGTGCTTAACTICTC i t f i e e c s h e i k r n h t n . r s i v r i s s f d c s n f i f . v f q i A FM L OS U KE 111 GATGGCGTGAGCTTAACCA CTACCGCACTCGAATTGGT	P2 PROMOTE AAAAGCAACGCAGTTTAG ITTITEGTTGCGTCAAATC k s n a v y k a t q f t c k q r s l l f l l a t . f a v c n v s f c r l k HH 8 NA 8 FE v 33 2 TATTCGGCCTGACAGAAG ATAAGCCGGACTGTCTTC f g l t e d y s a . q k	EXTENTAL EXAMPLE AND A BE E CO 3 21	TTTACCTGGCC m d p : c w t r n g p v f i s g f b v r f p g ! D H P S N E 1 1 CATCCCTTAA TTACAGGGAATT/	AGTAATIGTIT s l t k h q k i n k i n k c t m l l MI BI OI I TACCGCATACG	TTGTAACCAAAATTI t l v l k h m f , i n i g f k V n t k i f c q n f m p k l D H D P B P R O H 1 11 ATCAGCGAAACGATG TAGTGGGTTTGGTAG i s e t i s e k r s	2600 7 7 8 7
GACGGAACTGCGGA d g t a e t e l r k r n c g s p v e s l v s s r r f q p GATCAATTACTGTG CTAGTTAATGACACC d g l u f n y c g	TAGTGGGTGAGGGTTAACTTCTC i t f i e e c s h e i k r n h t n . r i v r i z s f d c s n f i f . v f q i A FM L OS U KE i 11 GATGGGGTGAGGTTAACCA CTACCGCACTCGAATTGGT d g v s i t i	P2 PROMOTE AAAAGCAAGGAGTTAAA k s n a v y k a t q f t k q r's l l f l l a t . f a v c n v s f c r l k HH	ERTEGETCACTEACT PATACTAAATTI y d k m i i i f k s h n l M MM A BB E CO 3 21 ACACACAGTEACC TOTGTCACTCACTCACTCACTCACTCACTCACTCACTCACTC	TTTACCTGGCC m d p : c w t r n g p v r i g g f h v r f p g i D H P S H E 1 1 TACAGGGAATT i s l i s p . j i l n	AGTAATIGTIT S	TTGTAACCAAAATTI t v k h w f . i n i g f k v n t k i f c q n f m p k D	- 2600 f

FIGURE 3I

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TTTTCAGAAAAACGATCGGCTETAAAGTGAATATCGAATGGGATATGATCGGAAAATATATGTATCGATTTTTGCATAAAGCCAATGAAAATAAGACCC
AAAAGTETTTTTTGCTAGCCGAGATTTCACTTATAGCTTACGCTATAGCTTTTATATACATAGCTAAAAACGTATTTCGGTTACTTTTATTCTGGG
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MCAMCCATTACAAAGCCTTCTTAAGCCATACGGCTTTTAGAGGGGACATTTGCATGCTTCATCCCGATAGAAGAAGCACTGGCCGCTTTAAAAAA
TTGTTTGGTAATGTTTTCGGAAGAATTCGCCTTTTGCCGAAAATCTCTCCTTCTAAACGTACAAAGTAGGCTATCTTCTTCTGTGACCTGCGAAATTTTTTT
 q titk a fla engf.rgrfacfir.kkhutl.kk
nkplqkps, aktafreedlh vaadrrst grfkk
tnhyksiikrkriierkicafh pieeaidaikk
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GGCGAAGTEATCATCGTTGTAGATGATGAAGAEAGAGAAAATGAAGGAGACTTTGTGGCTCTTGCCCAGCATGCAACGCCCGAAGTCATTAACTTTATGG
3000
CCGETTCAGTAGTAGCAACATCTACTACTTCTGTCTCTTTTACTTCCTCTGAAACACCGAGAACGGCTCGTACGTTGCGGCCTTCAGTAATTGAAATACC
aksssi.mmkteknketluilpsmqrrksitiu
rshhrer..rqrk.rrlegseraenagsh.lyg
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afdddnyiifvsfifsvkhakglmerridnykhirt..rqthbleifhtlsqpaqrahtapt..s.p
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FIGURE 3J

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GCTGTGTACCCTCTC	CTGACTAGACGT	GCGGCGAGTCACTCCTT1	AGDETCTGTCCGAACTAGAAG	TGGGATACCAACTCGTA	TATOTCTCAGAGTGGT	
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CACTGCATTTACCGT	AAGCATAGACCA	TCGTGAAACGAAGACAG	TATCAGCGCTCAAGAAAGAT			- 3200
GTGACGTAAATGGC	ATTCOTATCTGGT	AGEACTITGETTETGTE	CATAGTEGEGAGTTETTTETA	<u>CAAAATGGCAAGTTCGTA</u>	ACCACCTGTCGTTTAG	G
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			TGCGAAAAAGGAGGTGTCCT			- 3300
CACGGTAGACTAAA	AGTEGEAGGEECE	CETETALAAAGGTGACTA	ACCETTTTTTCCTCCACAGG	CTTTTCGCGCCCGGTAT	ETETTEGACGACAACT/	/C
al.f	4 2 4 9 1 q r p q	ehfstd hifpli	rkkevs.cekrop	e r r g p y k s 'a g h t	v d	΄.
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FIGURE 3K

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qqikriellrr, khf ahlrfapalaqa, qfqs
rasaqpdgpaptmqsiifsspvialtgssmsiaf
     M SM BND
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AAAGCATCAATTAAAAATGATCACCATTAAGGATTTGATTCAATACCGTTACAATCTGACAACATTGTCGAGGGTGAAGTTGACATTACGGTGCCTACT
                             --.-------. 3500
THTCGTAGTTAATTTTTACTAGTCGTAATTCCTAACTAAGTTATGCCAATGTTAGACTGTTGTGAACAGCTCGCACTTCAACTGTAATGCCGACGGATGA
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                                                     222
GATTTTGGGACATTTAAGGTTTATGGATACACAAATGAGGTAGATGGAAAGAGCATGTCGCATTTGTGATGGGAGATGTGCCGTTCGGAGAAGAACCGG
CTAAAACEETGTAAATTECAAATAECTATGTGTTTACTCCATCTACCTTTTCTCGTACAGCGTAAACACTACCCTCTACACGGCAAGCCTCTTCTTGGCC
ilghlrfædt qør. æksmeht.
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fwdi.giwihk.grwkracriedgreavertg
dfgtfkvygytnevdgkehvafvægdvpfgeepv
ikpekinisveilyistimdek.hhsihresttr
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FIGURE 3L

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H AA HHS
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 B VS PCC
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                                                   HBRSH1
 O AU AIR
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 2 21 211
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TATTGGTCCGGGTGCATTCAGAATGTCTCACAGGTGACGTGTTTGGCTCTCATCGCTGTGATTGCCGACCGCAGCTGCACGCCGCGCTGACCAAATTGC
ws g c i q n v s g v t c l a l i a v i a d r s c t p r . t k i p
igpgafrmshr.rvwlssl.trtaasrraepne
tvrvhsecttgdvfgshrcdcgpqthaatnqta
y q d p h m . f t e c t v h k a r m a t i e s r l q v g r q v l n
ipgpantid. thrtqsedshnrvasarresgfqtntrtceshrvpstnpe.rqsqpgcscsesfwis
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                  N A
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3800
GCGTCTTCCGGGCACCTCACGAGGACATGAACGCGGGTTCTTCCTGCGTAGCCAAATTAGTTATTTAATTTCGAATATTCGAAGTCCTTGTTCCGATA
 qkavecsctcakkdeesv.sin.klisfrnkam
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egrgvtlytrqegrgigtinktkayktqeqgy
g cfatsheq v q alfs sadt. dif. fs ilk i flai
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GACACCGTAGAAGCCAATGAGGCGCTTGGATTCTTGCCGGATCTTCGCAACTATGGCATCGGAGCACAAATTTTACGCGACCTCGGTGTCCGGAATATGA
CTGTGGCATCTTCGGTTACTCCGCGAACCTAAGAACGGGCTAGAAGCGTTGATACCGTAGCCTCGTGTTTAAAATGCGCTGGAGCCAAGGCCTTATACT
tp.kpmrridscrifatmasehkfyatsvsgi.
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FIGURE 3M

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FIGURE 3N

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	. c		AR YR N A 42 11 P 1	¥ 1	U K	0 A 1 H	
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•	GCGTAGACACA	MATGACATTGATGTGGC	•	TTTGAAATA	CCGTTTECTGCGAAAA	MATGGEGGAMCAMAMA	
						,	4300
'CGAGTCTGTAC	CGCATCTGTGT	TTACTGTAACTACACCG	MCCCMGGTCCGCGT.	MACTITAT	GGCAAACGACGCTTTTT	TTACCGCCTTTGTTTTTTT	
agtw	rrhi	. h e a	Lagrri	. n t	v	nggnkki	
						maetkk	
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						1 a s v f f	
						fhrfaff	
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		PL N.		V	w	ANA A U	
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TATGATGCTAT	TATCACATTE	GCACTGTCATCAGAGGC	GCAACGAEAEATTACG	ATTATGTET	GEAATGAAGETGEAAA	GGCATCGCGCAAGCAGCAA	
	•••.			•••••		,	4400
ATACTACGATA	ATAGTGTAACC	EGTGACAGTAGTCTCCG	CGTTGCTGTGTAATGC	TAATACAGA	CETTACTTCGACGTTTT	CCGTAGCGCGTTCGTCGTT	
	v h i a	hcharr	ndtle	1 . 1	a c k :	. h r a s s k	
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V	A	<u> </u>	9	Q	RAE		
1	1	3	1	1	221		
ACACTACTEGT	GTACCTGTCAT	CTTTGGAATTGTAACAA	TEAMACATCGAACA	GGCTATCGA	GCGTGCCGGCACAAAG	CGGGCAACAAAGGTGTAGA	
						,	4500
TGTGATGACCA	CATGGACAGTA	GAAACETTAACATTGTT	EACTTTTGTAGETTGT	CCGATAGCT	CCCACCCCCCGTGTTTTC	GCCCGTTGTTTCCACATCT	
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FIGURE 30

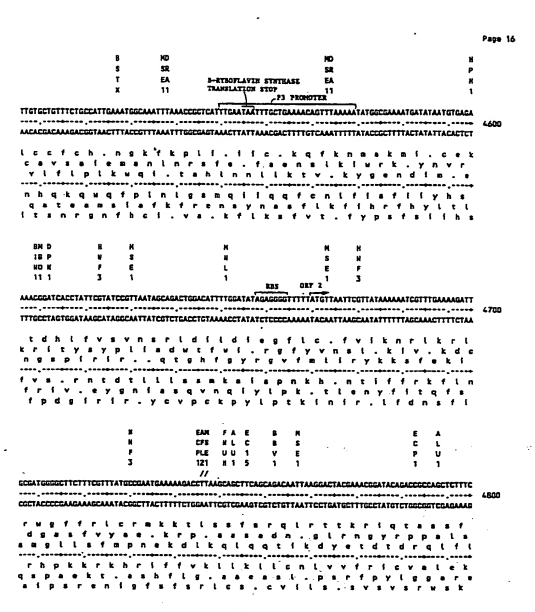


FIGURE 3P

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Page 17
                                        XND HB
    N
                       N
                         A
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                                        HBP P[
                         Q
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                                        211 21
 TYTGGAAAGAGGACGAGGATATCGTCGGAGCAATCGGAGTCGAAAAAAAGGATTCTGAGGTTGAGATCCGGCATATCAGTGTGAATCCTTCTCATCGCCA
 AAACETTTETCETGCTCCTATAGCAGCCTCGTTAGCCTCAGGCTTTTTTTTCCTAAGACTCCAACTCTAGGCCGTATAGTCACACTTAGGAAGAGTAGCGGT
 fgkrtrisse, qseskkrilrlrsgisv. illiai
lergrgyrrsnrsrkkgf.g.dpayqcesfssp
  tergrgyrrsnrsrkkgf, g.dpayq
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 k p f t v t i d d s c d s d f t i r t n t d p m d t h i r r m a
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TCAAGGAATCGGAAAACAGATGATGGATGCTTTAAAGCATTTATTCAAAACGCAAGTACTGGTTCCAAATGAATTAACGCAGAGCTTTTTCGAACGTTGT
AGTTCCTTAGCCTTTTGTCTACTACCTACGAAATTTCGTAAATAAGTTTTGCGTTCATGACCAAGGTTTACTTAATTGCGTCTCGAAAAAGCTTGCAACA
 kesenr.uml.siyskrkyufqmn
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 q g i g k q m m d s l k h l f k t q v l v p n e l t q s f f e r c
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             STOP
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GTTCCAGTCGTCGTAGTTCTGTAAAGTATGTTATTAATTCGTCTCCGACACTAGTCAGAGACGAAAAAAAGACGCAAGATAAAGAAAAAGTGCAAGTGC
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FIGURE 3Q

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Page 18
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                                                          113
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CTACTGCAGTCAGGCTAGGGCGTTTGCCACLAACAGCTATTCTTTATACAACGACTCACGTGACCCCACGGGGGTACATATGAAAAAAAGGACGTAAGC
ddvspipqtvfvdkkyvaectglppcilffpafd
atsvesekecisienalisaigaphvyffilhs
.eqsdpangvar.elaa.vhwaapmytfisaie
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ATCCTGCATGCTTCCTCCAGTTTCTCATCTTTGATTGGCAGTATAATGCTTTTATAGGCAGGAGGGTTTCGATTTGTTCGTAAACCGATTGCATAAGTT
TAGGAGGTAGGAAGGAGGTCAAAGAGTAGAACTAACCGTCATATTAGGAAAATATCGGTCTCTCGCCAAAGCTAACCAAGCATTTGGCTAACGTATTCAA
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COAGCAMACOGCCATGATCAAGCCCTAMGTCTTCGACTGCCCGGTGTTCTGCTTGAAGAATCCCGGATGCTGTTCGCCCATCAGTCTTTTTTGCCCCGGCTGT
GCTCGTTTGCCGGTACTAGTTCGGGATTCAGAAGCTGACGGCCACAAGACGAACTTCTTAGGCCTACGACAAGCGGTAGTCAGAAAAACGGGGCCCGACA
 eqtamikp.vfdcpvfclknpdavrhqsfcpgc
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FIGURE 3R

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reve	GAAGAEACTACTATATT	regetgaegttegaettattteggg	TGGGTTATCGEAAAGCA	MEMM	CCCCTAGAA	GGLAGG	TTATAAGAAG	A
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, q	rrhhyl kqssi MB LS AM	avalqifg fgscasylg H L A	v w y r k r g l l t k t H A	k k	D D d	e k i		r
, q	rrhhyl) kqssi _l MB LS	avalqifg fgscasylg N	v w y r k r g l l t k t H A I L	k k	D D d	e k i		r
, q	rrhhyl kqssii MB LS AM 31	avalqifg fgscasylg H L A 3	V W Y P K P G L L T K T H A I L H U 3 1	k k	D D d	e k i		r
, q	rrhhyl kqssii MB LS AM 31	avalqifg fgscasylg H L A	V W Y F K F G L L T K T H A I L H U 3 1	k k	D D d	e k 1 g e		r
TITA	HB LS AH 31 CATGGGATTCAAAATAAT	avalqifg fgscasylg N L A 3	V W Y P K P g l l t k t H A I L H U 3 1	k k	D D d	rgekı	4 y e e	·
TITA	HB LS AH 31 CATGGGATTCAAAATAAT	avalqifg fgscasylg H L A 3	V W Y P K P g l l t k t H A I L H U 3 1	k k	D D d	rgekı	4 y e e	·
T T T AAATT	FFR hyl NB LS AM 31 CATGGCATTCAAAATAAT GTACCGTAAGTTTTATTA	R CECUTOTCGCATGAAACTCAACGAG	H A I L H U 3 1	k k	D D d	rgekı	4 y e e	r
TATTT	HB LS AH 31 CATGGGATTCAAAATAAT	R C B S Y L G N L A 3 CCCCGTGTCGCCATGAAACTCAACGAG GGGCAACGCGTACTTTGAGTTGCTC	W Y P K P G L L T K T L H U 3 1 ATAATCTATATATATCTT	k k	D D d	eki ge	4 y e e	r
T T T AAATT	HB LS AN 31 CATGGCATTCAAAATAAT GTACCGTAAGTTTTATTA	BY BLQ IF 9 F 9 E C B S Y L 9 N L A 3 CCCCGTGTCGCGATGAAACTCAACGAG GGGCAACAGCGTACTTTCAGTTGCTC P C F B k l n e	W Y P K P G I I T K T I L H U 3 1 I L H U 3 1 I L H U 3 1 I L H U 3 1 I L H U	k k	D D d	rgekı ge	4 y e e	r
T T T AAATT	PP h h y l k q s s i i NB LS AN 31 CATGGCATTCAAAATAAT GTACCGTAAGTTTTATTA m a f k i i w h s k s h g i q n n	BY BLQ IF 9 F 9 E C B S Y L 9 N L A 3 CCCCGTGTCGCCATGAAACTCAACGAG GGGCAACAGCGTACTTTCAGTTCCTC P C F B k l n e F Y 4 . n s t F P Y S h e t q F d	W Y P K P G L L T K T L H U 3 1 EATAATCTATATATATCTATATATCTATATATCT	k k	D D d	rgeki	4 y e e	·
TITA	PP h h y l k q s s i i NB LS AN 31 CATGGCATTCAAAATAAT GTACCGTAAGTTTTATTA m a f k i i w h s k s h g i q n n	BY BLQ IF 9 F 9 E C B S Y L 9 N L A 3 CCCCGTGTCGCCATGAAACTCAACGAG GGGCAACAGCGTACTTTCAGTTCCTC P C F B k l n e F Y 4 . n s t F P Y S h e t q F d	W Y P K P G I I T K T I L H U 3 1 I L H U 3 1 I L H U 3 1 I L H U 3 1 I L H U	k k	D D d	rgeki	4 y e e	r

FIGURE 3S

FIGURE 4.

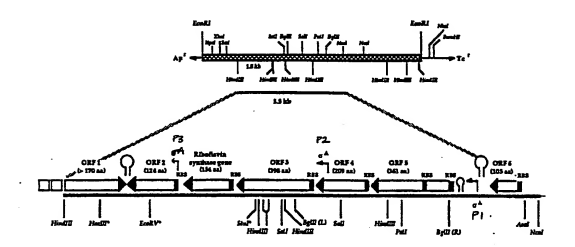


FIGURE 5.

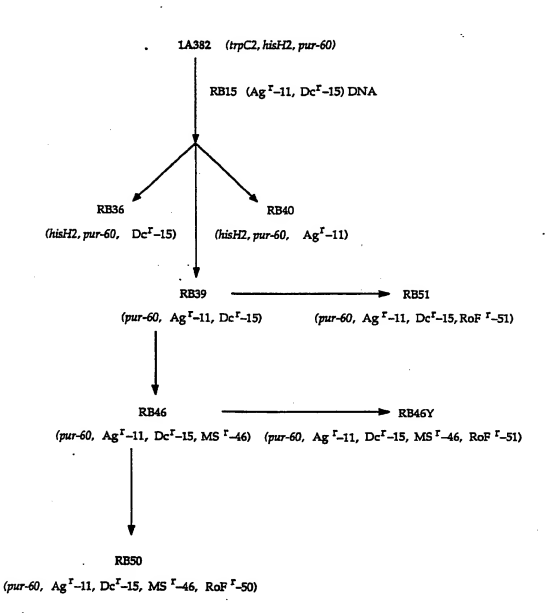


FIGURE 6.

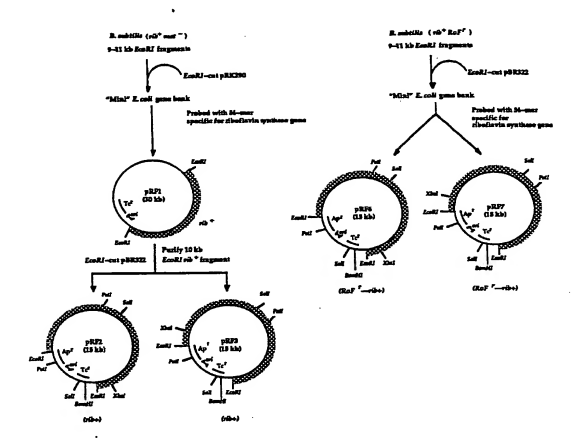


FIGURE 7.

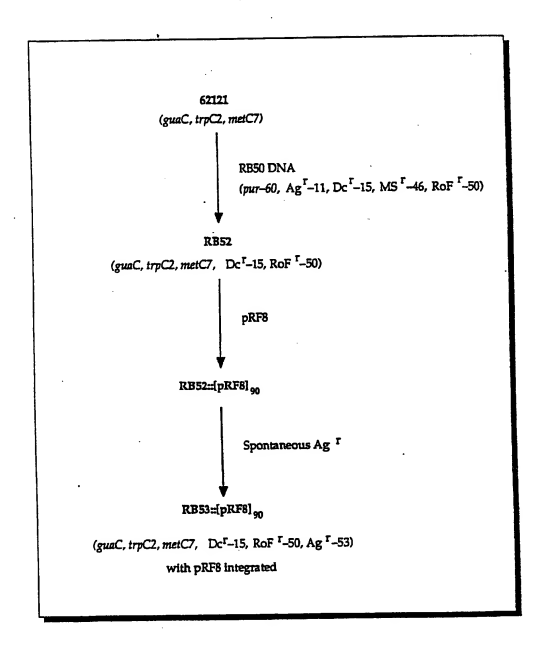


FIGURE 8.

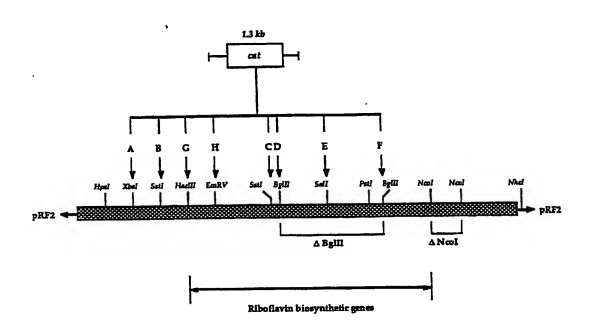
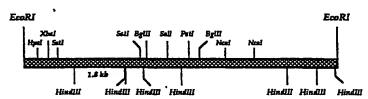


FIGURE 9.

<i>Upstream from</i> ribP1	Within 5' leader mRNA	At 3' end of rib operon
A G · C C · G T · A T · A T · G C · G A · T C · G A · T A · T A · T A · T	T T A T A T · A T · A T · A A · T G · C C · G C	G A T G•C T•A C•G G•T G•C A•T C•G G•C A•T C•G G•C A•T C•G G•C A•T
#708 #742	#1034 #1062	#5037 #5064
$\Delta G = -20 \ kcal/mol$	ΔG = –26 kcal/mol	$\Delta G = -16.5 \ kcal/mol$

FIGURE 10.



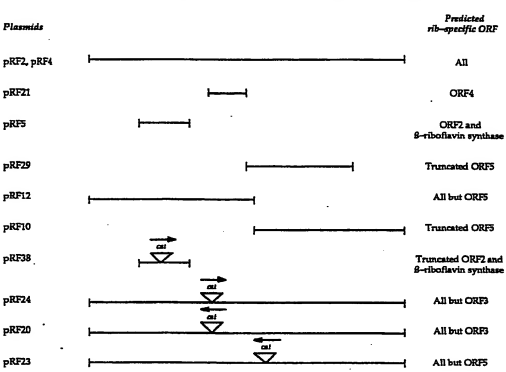


FIGURE 11.

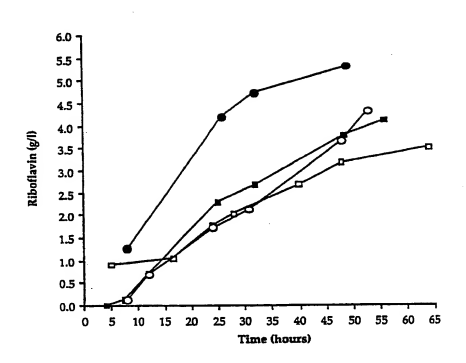


FIGURE 12.

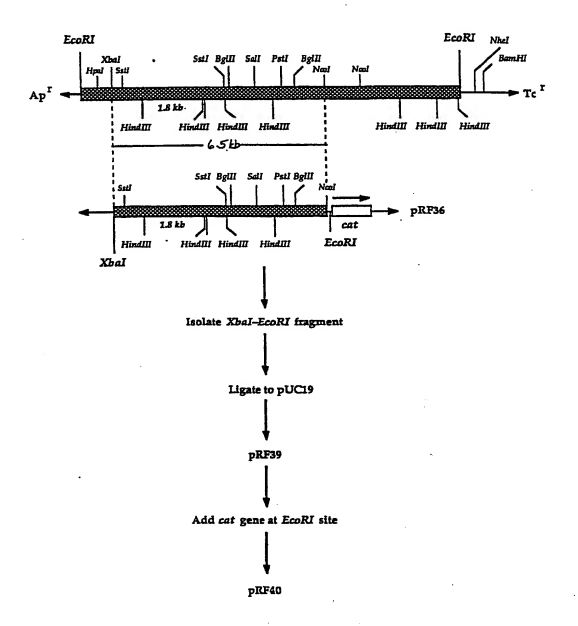


FIGURE 13.

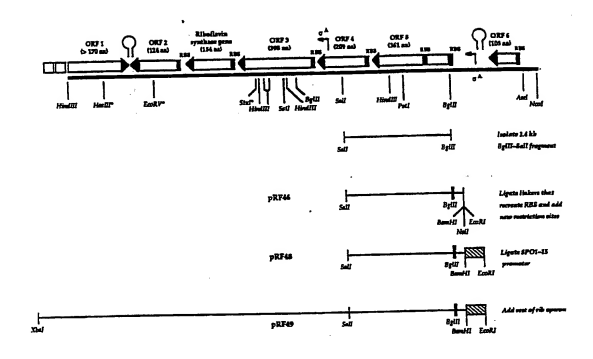


FIGURE 14.

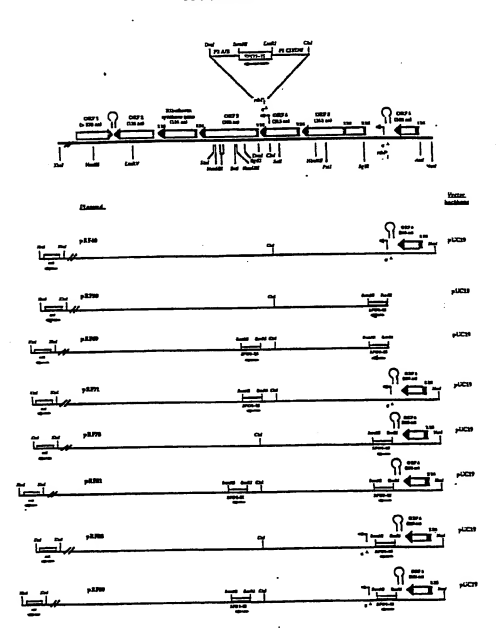


FIGURE 15.

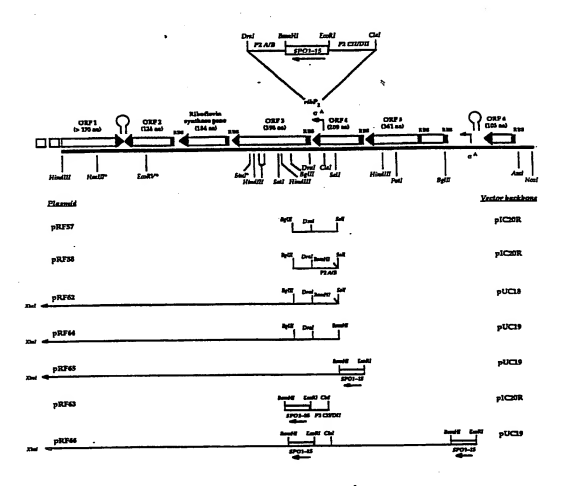
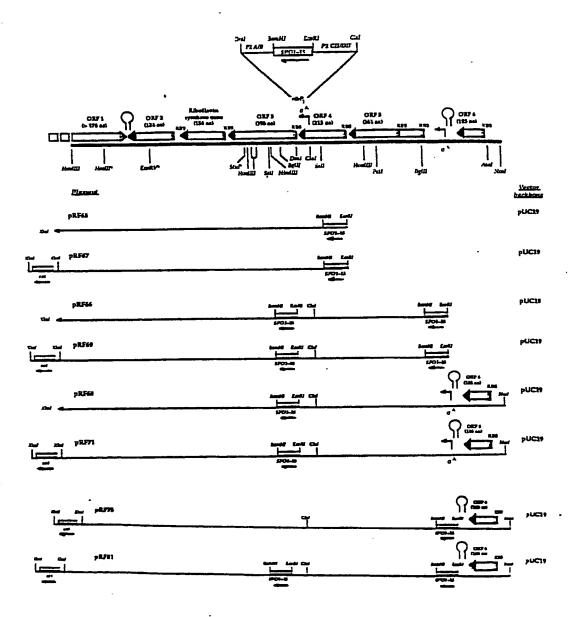


FIG. 16



F16.17

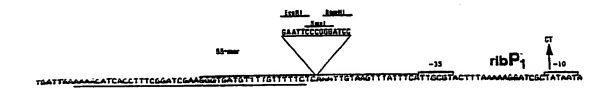


FIGURE 18

RB-5	AATTCATGCATGGATCCGACGGTAAATAAC AAAAGAGGGGAGGG
RB-6	GATCTAAGGCCAGCTTCATATAATACTCTT CCATTTGTTTCCCTCCCTCTTTTTGTTATT TACCGTCGGATCCATGCATG
P2-A	TCGACGGATCCTTTTAGAGAGGAAGATTTG CATGTTTCATCCGATAGAAGAAGCACTGGA CGCTTT
P2-B	AAAGCGTCCAGTGCTTCTTCTATCGGATGA AACATGCAAATCTTCCTCTCTAAAAGGATC CG
P2-CII	CGATTTTTGCATAAAGCCAATGAAAATAAG ACCCAACAAACCATTACAAAAGCCTTCTTA AGCGAAAACGGCTTTTAG
P2-DII	AATTCTAAAAGCCGTTTTCGCTTAAGAAGG CTTTTGTAATGGTTTGTTGGGTCTTATTTT CATTGGCTTTATGCAAAAAT

EUROPEAN SEARCH REPORT

EP 90 11 1916

	DOCCMENTS CONSI	DERED TO BE RELEVA		
Category	Citation of document with it of relevant pa	ndication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.5)
D,Y	FR-A-2 546 907 (V NAUCHNO-ISSLEDOVATE GENETIKI I SELEKTSI MIKROORGANIZMOV (VN * Whole document *	LSKY INSTITUT I PROMYSHLENNYKH	1-5,8- 13,15	C 12 N 15/52 C 12 P 25/00 C 12 N 1/20 / (C 12 N 1/20 C 12 R 1:125
Y	CHEMICAL ABSTRACTS, 5th June 1978, page 166562b, Columbus, RABINOVICH et al.: Escherichia coli ce subtilis riboflavin plasmids pPR1 and p NAUK SSSR 1978, 238 * Whole abstract *	293, abstract no. Ohio, US; P.M. "Expression in lls of the Bacillus operon in hybrid PR2", & DOKL. AKAD.	1-5,8- 13,15	C 12 R 1:19)
Y	CHEMICAL ABSTRACTS, 13th September 1982 no. 86284h, Columbu JOMANTIS et al.: "M successive cloning Bacillus chromosome NAUK SSSR 1982, 264 * Whole abstract *	, page 214, abstract s, Ohio, US; J. ethod for the of linked sites of s", & DOKL. AKAD.	1-5,8- 13,15	TECHNICAL FIELDS SEARCHED (Int. CL.5) C 12 N C 12 P
Y	PANINA et al.: "Clo riboflavin biosynth Bacillus subtilis o	ge 177, abstract no. Ohio, US; L.I. ning genes of the esis operon of n the plasmid pBR322 ia coli", & GENETIKA	1-5,8-13,15	
	The present search report has b	een drawn up for all claims		
	Place of search	Date of completion of the neuro		Examiner
TH	E HAGUE	28-09-1990	DE	SCAMPS J.A.
X : par Y : par do:	CATEGORY OF CITED DOCUME ricularly relevant if taken alone ricularly relevant if combined with an exmest of the same estegory hadogical background	E - carlles note	rinciple underlying on document, but point in the application of the document in the application of the reason	ublished on, or

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EUROPEAN SEARCH REPORT

Application Number

EP 90 11 1916

		DERED TO BE RELEVA			
Category	Citation of document with in of relevant par	dication, where appropriate,	t	elevant o ciaim	CLASSIFICATION OF THE APPLICATION (Int. CL5)
Y	CHEMICAL ABSTRACTS, 9th April 1984, pag 115822v, Columbus, RABINOVICH et al.: material in Bacillu (MOSCOW) 1984, 18(1 * Whole abstract *	e 138, abstract no. Dhio, US; P.M. "Cloning genetic s", & MOL. BIOL.		5,8- ,15	
Υ .	CHEMICAL ABSTRACTS, 5th November 1984, no. 164714r, Columb OKUNEV et al.: "Clo coli of a DNA fragm subtilis containing lysine and riboflav GENETIKA (MOSCOW), * Whole abstract *	page 157, abstract us, Ohio, US; O.V. ning in Escherichia ent of Bacillus the genes for in biosynthesis", &		-5,8- 3,15	•
Y .	CHEMICAL ABSTRACTS, 10th June 1985, pag 198921s, Columbus, RABINOVICH et al.: material in Bacilli BACILLI [PROC. INT. (Pub. 1984), 297-30 * Whole abstract *	e 160, abstract no. Ohio, US; P.M. "Cloning of genetic ", & BIOTECHNOL. CONF.]. 2nd 1983		5,8- 3,15	TECHNICAL FIELDS SEARCHED (lat. CL5)
	The present search report has a	een drawn up for all claims	\dashv		
	Place of search	Date of completion of the search		Τ	Examinar
TH	E HAGUE	28-09-1990		DES	CAMPS J.A.
Y: pa do A: tec	CATEGORY OF CITED DOCUME riferiarly relevant if taken alone riferiarly relevant if combined with an cument of the same category shoological background no-written discourse	E : earlier pais after the C	est docume ling date cited in th	ent, but pub e applicatio	ilshed on, or n